

Modeling the Performance and Cost of Lithium-Ion Batteries for Electric-Drive Vehicles

SECOND EDITION

Chemical Sciences and Engineering Division

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by

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ABBREVIATIONS

ASI	area specific impedance
BOL	beginning of life
DMC	dimethyl carbonate
EC	ethylene carbonate
EG/W	ethylene-glycol/water heat transfer fluid
EMC	ethyl methyl carbonate
EOL	end of life
EV	electric vehicle
Gr	graphite
GSA	General, Sales, and Administration
HEV	hybrid electric vehicle
HEV-HP	high-power assist hybrid electric vehicle
LCO	lithium cobalt oxide
LFP	lithium iron phosphate
Li	lithium
Li-ion	lithium-ion
LMO	lithium manganese spinel
LMR	lithium and manganese rich
LTO	lithium titanate spinel
microHEV	micro or mild power assist hybrid electric vehicle
MW	molecular weight
NCA	lithium nickel cobalt aluminum oxide

NMC	lithium nickel manganese cobalt oxide
NMP	N-Methyl-2-pyrrolidone
OCV	open-circuit voltage
OEM	original equipment manufacturer
PE	polyethylene
PET	polyethylene terephthalate
PHEV	plug-in hybrid electric vehicle
PP	polypropylene
R&D	research and development
SOC	state of charge
USGS	United States Geological Survey

LIST OF SYMBOLS

Section 3, 4 and 6

а	ratio of interfacial area to electrode volume, cm ⁻¹
A_{pos}	area of the positive electrode, cm ²
A _{term}	area of the terminal, cm^2
ASI _{energy}	area specific impedance for energy, ohm cm ²
ASIpower	area specific impedance for power, ohm cm ²
С	cell capacity, Ah.
C_1	parameter
C_p	specific heat capacity, J/g K
d_H	hydraulic radius, cm
е	battery to wheels efficiency
E	total energy, Wh
Ė	energy usage rate, Wh/mile
<i>f</i> _{Pd}	vehicle drag factor, kW/mph ³
f_{Pf}	rolling friction factor, kW/mph
F	Faraday constant, 96485.3 C/mol
G	mass flowrate, g/s
h	heat transfer coefficient, W/cm ² K
H_j	height of <i>j</i> , cm
H/W	aspect ratio of pouch cell
<i>i</i> _o	exchange current density related to the interfacial area, A/cm^2
Ι	average current density, A/cm ²
I_{lim}^{ionic}	ionic limiting current density, A/cm ²

In	local current density, A/cm ²
I _{total}	total current density, A
k	thermal conductivity, W/cm K
l_j	length of $_j$, cm
L_j	thickness of <i>j</i> , cm
m_j	mass of <i>j</i> , g
n	parameter
N_j	number of <i>j</i>
[N/P]	negative to positive capacity ratio
Р	battery power, W
P _{batt}	maximum designed battery power (rated power), W
P_a	accessory power, kW
P_d	aerodynamic drag power, kW
P_f	rolling friction power, kW
P_s	battery power to sustain speed S, kW
q	heating rate, W
Q	specific capacity of the electrode, mAh/g
r _C	C-rate, h ⁻¹
r _{C,lim}	limiting C-rate, h ⁻¹
<i>r</i> _j	radius of <i>j</i> , cm
\overline{R}	universal gas constant, 8.3144 J/mol K
R_j	resistance of <i>j</i> , ohm
S	vehicle speed, mph

t	time, s
Т	temperature, K
$U_{ocv,P}$	open-circuit voltage at SOC for power, V
$U_{ocv,E}$	open-circuit voltage at SOC for energy, V
ν	square root of dimensionless exchange current
V_{cell}	cell voltage, V
[V/U]	fraction of the open-circuit voltage
W_j	width of <i>j</i>
X	Cartesian coordinate, cm
X_{comp}	compression factor
α	constant, ohm cm ³
β	constant, ohm cm ²
Eact	volume fraction of active material
η^{1st}	first cycle efficiency
μ	fluid viscosity, g/cm s
$\Phi_{1,k}$	metal potential of foil k, V
$ ho_j$	density of j , g/cm ³
σ_j	conductivity of <i>j</i> , S/cm
Section 5.1	

С	final cost of lithiated oxide, \$/kg			
C_i	cost of raw material for component i, \$/kg			
C_o	baseline cost of lithiated oxide, \$/kg			
x_i	molar stoichiometry of component i			

MW_i	molecular weight of component i, g/mol
MW	molecular weight of lithiated compound, g/mol
Section 5.4 C	capital cost of an installed equipment for the designed battery, \$
C_o	capital cost of an installed equipment for the baseline plant battery, \$
р	power factor
R	designed battery processing rate for specific process step
R_o	baseline plant processing rate for specific process step

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PREFACE TO THE SECOND EDITION

The BatPaC model and report were first openly distributed to the public, free of charge, in November 2011 from the website www.cse.anl.gov/batpac. During the first year of release, over 450 independent downloads have occurred from leading universities, laboratories and companies around the world. Users of the model include PhD students, policy makers, high-profile startups, and Fortune 500 corporations. Some have modified the model for their own use, while others use it as is.

The major updates included in BatPaC v2.1 include the following:

- Air-based thermal management option
- Automatic calculation of uncertainty in point cost estimate
- Updated transition metal based cathode material costs
- Parallel pack and module option
- Improved heat generation estimation
- Multiple cost-breakdown formats

Currently, the calculations are based in a Microsoft[®] Office Excel spreadsheet. Instructions are provided for use; however, the format is admittedly not user-friendly. A parallel development effort has created an alternate version, the Battery Production Model (BatPro), based on a graphical user-interface that will be more intuitive to some users. BatPro is available from the Argonne software shop for a nominal fee.

EXECUTIVE SUMMARY

This report details the Battery Performance and Cost model (BatPaC) developed at Argonne National Laboratory for lithium-ion battery packs used in automotive transportation. The model designs the battery for a specified power, energy, and type of vehicle battery. The cost of the designed battery is then calculated by accounting for every step in the lithium-ion battery manufacturing process. The assumed annual production level directly affects each process step. The total cost to the original equipment manufacturer calculated by the model includes the materials, manufacturing, and warranty costs for a battery produced in the year 2020 (in 2010 US\$). At the time this report is written, this calculation is the only publically available model that performs a bottom-up lithium-ion battery design and cost calculation. Both the model and the report have been publically peer-reviewed by battery experts assembled by the U.S. Environmental Protection Agency. This report and accompanying model include changes made in response to the comments received during the peer-review.

The purpose of the report is to document the equations and assumptions from which the model has been created. A user of the model will be able to recreate the calculations and perhaps more importantly, understand the driving forces for the results. Instructions for use and an illustration of model results are also presented. Almost every variable in the calculation may be changed by the user to represent a system different from the default values pre-entered into the program.

The distinct advantage of using a bottom-up cost and design model is that the entire power-toenergy space may be traversed to examine the correlation between performance and cost. The BatPaC model accounts for the physical limitations of the electrochemical processes within the battery. Thus, unrealistic designs are penalized in energy density and cost, unlike cost models based on linear extrapolations. Additionally, the consequences on cost and energy density from changes in cell capacity, parallel cell groups, and manufacturing capabilities are easily assessed with the model. New proposed materials may also be examined to translate bench-scale values to the design of full-scale battery packs providing realistic energy densities and prices to the original equipment manufacturer.

1. Introduction

The recent penetration of lithium-ion (Li-ion) batteries into the vehicle market has prompted interest in projecting and understanding the costs of this family of chemistries being used to electrify the automotive powertrain. The model described here-in is a calculation method that was developed at Argonne National Laboratory (Argonne) for estimating the manufacturing cost and performance of Li-ion batteries for electric-drive vehicles including hybrid-electrics (HEV), plug-in hybrids (PHEV), and pure electrics (EV). To date, a number of cost models of various levels of detail have been published in different forms.¹⁻¹¹ The cost of a battery will change depending upon the materials chemistry, battery design, and manufacturing process.¹²⁻¹⁴ Therefore, it is necessary to account for all three areas with a bottom-up cost model. Other bottom-up cost models exist but are not generally available and have not been explicitly detailed in a public document. The motivation for this work is based on a need for a battery cost model that meets the following requirements:

- 1. Open and available to the entire community
- 2. Transparent in the assumptions made and method of calculation
- 3. Capable of designing a battery specifically for the requirements of an application
- 4. Accounts for the physical limitations that govern battery performance
- 5. Based on a bottom-up calculation approach to account for every cost factor

The Battery Performance and Cost model (BatPaC) described here-in is the product of long-term research and development at Argonne. Over a period of years, Argonne has developed methods to design Li-ion batteries for electric-drive vehicles based on modeling with Microsoft[®] Office Excel spreadsheets.¹²⁻²⁰ These design models provided all the data needed to estimate the annual materials requirements for manufacturing the batteries being designed. This facilitated the next step, which was to extend the effort to include modeling of the manufacturing costs of the batteries. In the following sections of this document, a model is presented that meets the above criteria and may be used to analyze the effect of battery design and materials properties on the cost of the final battery pack. Use of BatPaC requires some basic knowledge of battery packs; however, a user does not need to be an expert. For instance, the number of cells and thus battery pack voltage must be specified by the user. However, default values are available for more specific requirements such as experimentally measured values. In this way, a person with reasonable knowledge of batteries may be able to conduct cost comparisons and "what if" studies.

The battery pack design and cost calculated in BatPaC represent projections of a 2020 production year and a specified level of annual battery production, 20,000-500,000. As the goal is to predict the future cost of manufacturing batteries, a mature manufacturing process is assumed. The model designs a manufacturing plant with the sole purpose of producing the battery being modeled. The assumed battery design and manufacturing facility are based on common practice today but also assume some problems have been solved to result in a more efficient production process and a more energy dense battery. Our proposed solutions do not have to be the same methods used in the future by industry. We simply assume the leading battery manufacturers, those having successful operations in the year 2020, will reach these ends by some means.

Establishing the validity of the model calculation is important in justifying the conclusions drawn from exercising the model. The report and model have been subjected to a public peerreview by battery experts assembled by the U.S. Environmental Protection Agency. Changes have been made in response to the comments received during the peer-review. The design methodology used has been previously validated against cylindrical wound cell formats.¹⁵ The calculated materials quantities agreed with the actual values within 3 %. Moving to a prismatic format simplifies the current collection calculation while leaving the governing equations unchanged. The new approach developed for calculating the cell impedance has been validated against experimental measurements from electrodes up to 100 µm in thickness.²⁰ The module and battery jacket construction is of lighter construction compared to contemporary designs. The battery pack energy density calculated in BatPaC is higher than the battery packs used in the first versions of the Nissan Leaf and Chevrolet Volt for equivalent cell designs (calculated value of 100 Wh/kg compared to a reported value near 84 Wh/kg for the Volt²¹). Significant engineering advances are necessary to minimize the current inactive material burden in the commercial pack designs, thereby reducing the cost, mass and dimensions of future automotive battery packs. We have assumed a design that we believe will be representative of the engineering progress achieved by successful manufacturers in the year 2020.

Validation of the input material and capital costs are more difficult to achieve as few values are publically available. We have relied, to a large extent, on private communications from equipment manufacturers, materials suppliers, cell manufacturers, and original equipment manufacturers (OEM). Variation does exist amongst the communicated values and we have maintained a practical level of skepticism for their accuracy. Experts from all aspects of battery development have reviewed the model both privately and as part of a formal peer-review process. While the largest uncertainty in calculated values will exist in point cost estimates, the most instructive information may be gained by examining ranges in parameter values and relative changes between material properties (*e.g.* the advantage of moving to a manganese spinel cathode from a layered-oxide material or from increases in cell capacity, etc). An initial variation study may be found in Appendix A.

The battery pack price to the OEM calculated by the model inherently assumes the existence of mature, high-volume manufacturing of Li-ion batteries for transportation applications. Therefore, the increased costs that current manufacturers face due to low scale of production, higher than expected cell failures in the field, and product launch issues are not accounted for in the calculation. The model results for year 2020 could be considered very optimistic if the transportation Li-ion market fails to develop as a result of insufficient investment in product research and development, reduced motivation for lowering petroleum consumption and greenhouse gas emissions, and/or a series of high-profile safety incidents.

2. Cell and Battery Pack Design Format

Various cell and battery design concepts are under development at battery manufacturers. Based upon experience gained from extensive previous work, we have found the exact design of the battery does not have an important effect on the cost for a set cell chemistry system; the amounts of electrode materials and the number, capacity and electrode area of the cells, are the determining cost factors. The most common cell designs for batteries nearing large-scale production are cylindrical wound cells, flat wound cells, and prismatic cells with flat plates. Cylindrical cells probably have a slight advantage for the assembly of the electrode-separator unit because of the ease of making a cylindrical winding. For the different cell designs, there are small differences in the weights of the terminal extensions and the procedures for connecting these extensions to the current collector sheets, with a small advantage for flat plate cells. The flat-wound and flat-plate cells form a more compact module and have better heat rejection capabilities than the cylindrical cells. These small differences would have minor effects on the cost of batteries produced in high volume in a mature, automated production plant and all of the cell designs can be adequately cooled for most applications. We conclude that the cost calculations would be relevant for batteries differing considerably from the selected design approach.

To provide a specific design for the calculations, a prismatic cell in a stiff-pouch container was selected. The terminals are almost as wide as the full width of the cell with the positive terminal at one end of the cell and the negative terminal at the opposite end. The electrical performance of a cell with this construction is near optimum, with a very low fraction of the resistance in the current collection structure. This cell design is amenable to either liquid or air based thermal management approaches. In this study, most of our attention has been directed to batteries using liquid thermal management. For that approach, the cells are enclosed in hermetically sealed modules, which are cooled on their exterior surfaces by ethylene glycol-water solution. The module enclosure protects the cell terminals from the heat transfer fluid, which is an electrolyte. As an additional safety precaution, a dielectric liquid could be used as the heat transfer fluid, such as a transformer coolant, but that approach has not been studied in the work of this report.

For air thermal management, the broad cell area must be individually contacted by the fluid due to the poorer heat transfer properties of air. Therefore, the cells are enclosed in an aluminum sleeve which provides air flow channels; the module enclosure is open at the bottom and top to accommodate upward flow of air through these channels. Both the liquid and the air thermal management designs are configured to be compact, light-weight and amenable to low-cost manufacture. It is unlikely that we have selected the most viable designs in this short study; there may be serious flaws in some details. However, the calculated overall performance and low cost for the selected design will be challenging to match in actual production and will only be met by the most successful manufacturers, those that will dominate the market.

The paragraphs below in section 2, provide the overall cell, module and battery pack design formats. Additional design details and methods of calculations are provided in sections 3 and 4.

2.1 Cell Design

The prismatic cell of this design embodies individual positive and negative electrodes consisting of current collector foils coated with electrode materials on both sides. The current collectors are usually solid copper and aluminum foils for the positive and the negative. An illustration of a segment of the cell is detailed in Figure 2.1. Each electrode is made up of active material particles held together by a polymeric binder. A conductive additive, carbon black and/or graphite, is added to the positive electrode and sometimes to the negative electrode. The electrodes and separator each have porosity that is filled with the electrolyte solution. During discharge, the Li-ions move from the electrode particles into the electrolyte, across the separator, and then insert into the particles composing the opposite electrode. The electrons simultaneously leave the cell through the current collection system and then enter through the opposite side after doing external work. The materials currently used in Li-ion cells are based on an intercalation process. In this process, the Li-ion is inserted into or removed from the crystal structure of the active material. The oxidation state of the active material, or host, is concurrently changed by gaining or losing an electron. Other electrode materials based on conversion reactions or electrodeposition could be implemented into the model if the user desired.



Figure 2.1. Cell sandwich inside of prismatic pouch cells.

The electrodes are easily and efficiently prepared by coating wide sheets of foil up to 2-meters in width with uncoated strips running the lengths of the foil being coated. The individual electrodes can be cut from these sheets with little waste of electrode coating material or foil (Fig. 2.2).

The separator for these cells can be handled as a single sheet that is folded back and forth as the electrodes are inserted. The electrodes are inserted so that all of the positive tabs extend beyond the separator sheet in one direction and the negative tabs extend in the opposite direction. The design model selects the number of electrodes to meet a set cell thickness determined by the type of cell: HEV, 6 mm; PHEV, 8 mm; EV, 12 mm. These cell thicknesses are default values and may be changed to suit the designer.

The stiff-pouch containment for the cell and the terminal seal is illustrated in Fig. 2.3. The cell terminals are formed from flat stock to be almost as wide as the entire cell. They are bent to the required shape and ultrasonically welded to the current collector tabs. The cell stack is then sealed between the two halves of the cell container. The cell housing material is a tri-layer consisting of an outer layer of polyethylene terephthalate (PEP) for strength, a middle layer of 0.1-mm aluminum for stiffness and impermeability to moisture and electrolyte solvent vapors and an inner layer of polypropylene (PP) for sealing by heating.^{18,19} The two halves of the cell container are pre-shaped to facilitate assembly. The aluminum foil in the cell container material provides stiffness and it may be increased in thickness to assist in conducting heat to the module container. After sealing the edges of the cell, the edges are flattened along the sides of the cell to form a compact shape. For liquid based thermal management, an aluminum conduction channel is added to assist in heat rejection at the sides of the cells (Fig. 2.3).



Figure 2.2. Illustration of coated current collector foil showing four rows of prismatic electrodes before slitting or stamping into individual electrodes

For batteries utilizing air based thermal management, the cell design format is the same as for liquid-based systems. To provide space for air flow between the cells, the cells are enclosed in an aluminum sleeve that provides the flow passages as illustrated in Fig. 2.4. The ridges between the air flow passages are 2-mm wide. The passages themselves are about 10-mm wide, the exact width being calculated to provide an integer number of passages across the length of the positive electrode.



Figure 2.3. Prismatic cell in stiff pouch container with aluminum conduction channel added for heat rejection from a liquid cooled module



Figure 2.4. Provision for cooling cells with cabin air

2.2 Module Design

2.2.1 Modules Using Liquid Thermal Management

The module format is based on a casing of 0.5-mm thick aluminum that is sealed by double seaming, a process that is well established and inexpensive because it is automated, rapid, and uses low-cost equipment that is common in the container industry. The sealing of the module provides an additional barrier to the loss of electrolyte solvent from the cells and the entrance of water vapor. These deleterious transfers through the seals of pouch cells may shorten their lives to less than the desired fifteen years.¹⁸

The cells are placed on their sides in the module and the terminals of adjacent cells are connected either mechanically with rivets and flat springs to maintain contact or by laser welding. Space is provided within the module casing on the left side, as sketched in Fig. 2.5, for an electronics package that includes cell monitoring for malfunctions (temperature and voltage) and for state-of-charge (SOC) control. The SOC control is activated whenever the battery is at rest and it diverts charge from the cells at highest voltage to those at lowest voltage.



Module with Sealed Aluminum Container Designed for Liquid Cooling

Figure 2.5. Module with hermetically sealed aluminum container for batteries utilizing a liquid thermal management approach

2.2.2 Modules Using Air Thermal Management

For systems using air as the heat transfer fluid, the module must be open to allow air to flow between the cells utilizing a larger surface area for heat transfer (Fig. 2.6).



Figure 2.6. Module using air heat transfer fluid with polymer container open at top and bottom

2.3 Battery Pack Design

2.3.1 Batteries Using Liquid Thermal Management

The model designs the battery pack (Fig 2.7) in sufficient detail to provide a good estimate of the total weight and volume of the pack and the dimensions of the battery jacket so that its cost can be estimated. The modules are arranged within the battery jacket either in a single row, with the terminals facing the same side of the pack, or in an even number of rows with the terminals in one row facing the terminal of an adjacent row. For a pack with a single row of modules, a busbar must be provided to carry the current to the front of the battery pack. This feature results in an additional cost for the busbar. For batteries with more than one row of modules (Fig. 2.7), the terminals are laid out in the module so as not to interfere with those on the opposite row of modules, thus conserving space in the battery pack. The modules in a row are interconnected, negative to positive terminals, by copper connectors. The modules casings are compressed together between steel sheets at each end of the battery pack. The compression force is applied by steel bands wrapped around the top and bottom of the row of modules. The compression is necessary to ensure intimate contact between the active layers that make up the pouch cells that are tightly fit into the modules. The compressive force also serves to add structural support to the module casings.

The modules are supported by a tray that positions the modules mid-way between the inside top and bottom of the pack jacket and provides space for heat transfer fluid to flow so that it cools or heats the top and bottom of each module. The rows of modules are assembled, compressed between end plates, interconnected and attached to the module tray. The module tray is attached to the pack closure at the front of the pack so that all connections to the pack terminals that lead to the exterior of the pack and signal wire feedthroughs can be made before inserting the attached modules into the jacket and making the final closure. The bolts depicted in the diagram (Fig 2.7) for making this closure are only illustrative. The battery jacket consists of a sheet of aluminum on each side of a 10-mm thick layer of ridged, light-weight high-efficiency insulation. The thickness of each of the aluminum layers is selected by the modeling program to be 1- to 2-mm thick, depending on the total volume of the modules. The insulation slows the interaction of the battery with the external environment that cools the battery in the winter and heats it in the hot summer weather.¹³



Pack with One Row of Modules

Figure 2.7. Insulated battery jacket with enclosed modules that are exposed on their upper and lower surfaces to an ethylene glycol-water heat transfer fluid.

Although the main purpose of the battery pack design for the model is to provide a plausible list of materials to estimate the manufacturing cost of the battery, the overall design approach permits the battery to be shaped by the designer to fit dimensional objectives. If there is a height restriction for the battery pack, a high ratio of length-to-width for the positive electrode will result in a battery of low height. Because the cell terminals are nearly as wide as the electrodes in the cell design selected for this model, the current collector structure adds very little to the total cell impedance even for electrodes with length-to-width ratios as high as 3.0. If the cell terminals were both brought out of the same end of the cell and, therefore had to be narrow, the resistance of the terminals and of the current collector foil in the vicinity of the terminals would be high, especially for long, narrow cells. Further discussion of the means of adjusting the shapes of the modules and the pack is provided in section 7.

2.3.1 Batteries Using Air Thermal Management

The air thermal management battery format is similar to the liquid thermal management format in that the cells are the same and the modules casings are held together by steel sheets at the ends of the row of modules and compressed by steel straps. The battery pack for air thermal management differs, however, in that much more space must be allotted for air flow than for liquid flow (Fig. 2.8).



Figure 2.8 Battery pack utilizing air thermal management approach.

Air is admitted along the entire length of the bottom of the pack to minimize the thickness of the flow passages above and beneath the modules. The air leaving the module is ducked to the inlet side of the fan, which can be located nearby to suit the vehicle design. The pack-long entrance and exit passages are insulated for thermal efficiency. These additions and the air passages between the cells and the wide passages required above and below the modules add considerable volume to air thermal management packs over that required by liquid thermal management packs. The thermal calculations described in section 4 indicate that cabin air-cooling is sufficient for HEV battery packs and for some larger battery packs, but it may limit the driving profiles that are feasible.

3. Modeling of Battery Design and Performance

The design portion of the model calculates the physical properties of a battery based on userdefined performance requirements and minimal experimental data. An illustration of the model is shown in Figure 3.1. The user is asked to enter a number of design parameters such as the battery power, number of cells and modules, and target voltage at maximum power, etc. In addition, the user must enter one of the following three measures of energy: battery pack energy, cell capacity, or vehicle electric range. Defining one of these values will determine the value of the other two. An iterative procedure then solves for the user defined energy parameter (energy, capacity, or range) and remaining battery properties by varying the cell capacity and electrode thickness. The result is the dimensions, mass, volume, and materials requirements for the cells, modules, and battery pack.

The model has been designed to allow the user to enter as many customized values as desired. In this way, the model allows flexibility in the battery chemistries studied and some of the cell, module, and battery design aspects. Hence, the focus of this report is on the method of calculation and not the exact values chosen for a specific capacity or cell thickness. However, the default cell design parameters as well as experimental data measured at Argonne National Laboratory, for a number of different battery chemistries both commercial and developed at Argonne, are available for use within the model. There are five governing equations for battery performance that calculate the current density, battery energy, electrode area, electrode thickness, and resistance. The voltage at maximum power and the area specific impedance (ASI) are two important parameters in the design model for calculating the battery performance. Most of the discussion will be spent on these two properties.

3.1 Criteria for Power, Energy, and Life

In order to fully specify a battery design, the user of BatPaC must supply criteria for power, energy, and life. These criteria will depend on the application for which the battery will be used. While the users may change some of the settings as they prefer, we list our suggestions in Table 3.1. The battery type is defined by the end-use application. Hybrid electric vehicles (HEVs), plug-in hybrid electric vehicles (PHEVs), and electric vehicles (EVs) have increasing levels of electrical energy storage for use by the vehicle drivetrain. The model will use Table 3.1 or the user's explicit inputs to size the battery correctly for the chosen application.

Battery Type	microHEV	HEV-HP	PHEV	EV				
SOC for Rated Power, %	50	50	20	20				
Power Duration, sec	2	10	10	10				
SOC Range for Useable Energy, %	40-65	40-65	20-90	10-95				
Cell Thickness, mm	6	6	8	12				

Table 3.1 Criteria for designing batteries for a specific end-use application

The microHEV is a micro or mild-hybrid that provides a moderate power level, ~ 25 kW, for two seconds. This design is best suited for cell chemistries capable of very high power-to-energy (P/E) ratios. The HEV-HP is a power-assist hybrid that provides the rated power for a full 10

second pulse. The power for both HEV applications is rated at 50 % state-of-charge (SOC). The energy available for discharge and charging is 25 % of the total energy to ensure long cycle life. As the capacity of the HEV cells is typically small, a cell thickness of 6 mm is used. The PHEV utilizes a much larger portion of the total energy, 70 %. At the end of discharge, the PHEV battery is operated in a charge sustaining mode. Therefore, the power rating for the battery is determined at 20 % SOC. PHEV cells should be much larger than HEV cells and thus a cell thickness of 8 mm is assumed. Finally, EV batteries use 85 % of their total energy with their power rated near the end of discharge. EV cell thicknesses are set to 12 mm.

Battery Design Model

Pack Requirements

- power
- energy or range
- number of cells

Key Constraints

- max electrode thickness
- target cell potential, V, at peak power
- assumed cell/module format

Iterative Spreadsheet

Solves for cell capacity and designs battery pack by varying:

- 1. Cell area
- 2. Electrode thickness
- 3. Internal resistance

Cell Chemistry

Measured Properties

- pulse power ASI
- sustained discharge ASI
- mAh/g, g/cm³
- electrode porosity
- SOC window
- physical properties
- ASI = area specific impedance

Figure 3.1 Summary flow of the design model

Calculated Battery Properties

- volume & mass
- specific energy, power
- materials required
As noted later in the report, selecting a parallel arrangement of cells automatically assumes a cell thickness of 6 mm regardless of the end-use application. Additionally, the use of negative electrodes operating at potentials high above the lithium metal potential may extend the upper end of the available SOC range from 95 to 100 %. The lithium titanate spinel, $Li_4Ti_5O_{12}$ (LTO), negative electrode is an example of an intercalation electrode with almost no risk of plating lithium metal during a charge pulse. On this basis, the available energy for LTO-based Li-ion cell is increased by 5 % to be 75 % for PHEVs and 90 % for EVs. In an established factory, the fixed design parameter is most likely the electrode area for a single layer rather than a set cell thickness. To make higher capacity cells, more layers of the predetermined footprint are stacked, thus increasing the cell thickness. In our model, the plant is constructed for the sole purpose of building the battery being designed. Flexibility to produce other products is not taken into account. Most importantly, the model calculates very little difference in cost whether the cell has large-area layers that are few in number as compared to a high number of layers. The area of each layer is set by the cell capacity requirement.

Accounting for capacity and power fade in the battery requires the user to design the battery with the appropriate excess energy and power at the beginning-of-life (BOL). Achieving rated end-oflife (EOL) power at a high fraction of the open-circuit voltage at BOL is one way to set the allowable power fade over the life of the battery. This is discussed in detail in the following section. Capacity or energy fade may be managed in multiple ways. The BOL SOC range for useable energy in Table 3.1 may be considered one approach to oversizing. As the battery capacity is reduced resulting from fade mechanisms, the loss in total capacity and energy is not as obvious when only cycling in the reduced SOC range. In addition, the SOC window may shift to different defined pack voltages to obtain a near constant useable energy value. Current practices by the OEMs suggest that the customer will accept some degree of capacity fade from the battery over the life of the vehicle. The Chevrolet Volt battery warranty states the capacity may fade by 10 - 30 % over the warranty period. Certainly, the customers' previous experience with consumer electronics devices has prepared them for a reduction in battery capacity with the life of the product. However, we do not hold the same opinion on power fade. We have allotted for significant power fade by designing EOL power to be achieved at a high fraction of the open circuit voltage at BOL. The design model does not attempt to predict fade rates or even suggest an allowable fade for a specific application, other than the SOC window for useable energy. It is our view that many aspects of materials chemistry, cell design, and battery use directly affect the decay rate of the battery pack. Hence, we allow the user to make any additional accommodations for decay believed to be necessary by entering larger total energy content than what is calculated from the useable SOC windows.

3.2 Voltage at Rated Power

The voltage at which a cell reaches the rated power is one of the most important factors in the design of a battery. However, this specification is one of the least discussed aspects of battery design. Our design approach assumes the drivetrain will never draw a power level greater than the rated power over the entire life of the battery. The voltage at rated power is a measure of the largest polarization the cell will undergo during operation at the BOL. This initial value has a

direct effect on round-trip battery efficiency, heat removal requirements, cold-cranking power, and allowable power fade. A basic calculation demonstrates the maximum achievable power for a battery at BOL is at 50 % of the open-circuit voltage (OCV). Operating at these conditions would result in an inefficient battery and require a significant cooling system to reject heat. More importantly, the battery will never be able to reach this power level after any increase in impedance occurs. With all certainty, the impedance of a battery will rise with time and the power rating of the battery will no longer be accurate.

We design the battery to achieve EOL power capabilities (rated power) at a specified fraction of the open-circuit voltage, [V/U], at BOL. This approach is unique when compared to current design practice of OEMs and cell manufacturers. However, a characteristic value of [V/U] exists for all batteries regardless of the battery design process. One may determine this value for an existing system in a straightforward manner. The voltage at rated power is measured at the end of a 10 s pulse at the EOL power rating and the SOC used for the power rating of a specific battery type (HEV, PHEV, EV). The designed [V/U] value is the measured potential at the end of the pulse divided by the open-circuit potential reached long after the pulse. This design point then captures the degree to which the battery has been oversized to enable long-life, cold-start, and efficient operation. The remainder of this section presents a discussion for setting the BOL voltage at rated power at no less than 80 % of the open-circuit voltage, [V/U] = 0.8. Defining the voltage as a fraction of the OCV, allows for direct calculation of all the necessary battery properties (see for example Eq. 3.6 or 3.8 in the section 3.3).

The allowable increase in battery resistance over the life of the battery is a function of the designed voltage for rated power. In general, designing the battery to achieve rated power at a higher [V/U] allows for larger resistance or impedance increases over the lifetime of the battery. Figure 3.2 created from Eq 3.1 displays how the voltage at rated power will change to meet the designed power as the internal resistance of the battery increases. Clearly, achieving BOL power at a high fraction of the OCV allows for greater degradation within the usable lifetime of a battery. R_1 is the initial resistance of the battery at BOL while R_2 is the resistance as the battery ages. If the minimum voltage is 55 % of the OCV, the allowable increase in resistance for batteries designed for BOL rated power at 70, 80, and 90 % OCV is 18, 55, and 175 %. The consequence of achieving the power at lower and lower fractions of the battery, Figure 3.2b and Figure 3.3. The proper design of a battery will account for the changes over the entire lifetime and not just desired behavior at BOL.

The level of heat production is significantly different at BOL for batteries designed to meet rated power at differing fractions of the open-circuit voltage. We may compare the differences in designed [V/U] by assuming the resistive heating (joule heating) is the most significant factor in determining the heat generation, Eq. 3.2. This assumption is true for moderate to high rate applications. We also reasonably assume the ASI will not change significantly in the range of current densities and electrode thicknesses we vary in the comparisons. We can analyze the difference in heat generation for different [V/U] values, Eq 3.3.



Figure 3.2 a) Required change in [V/U] to maintain rated power with increases in internal resistance over the life of the battery. b) Increase in current due to lowered [V/U].

$$\left[\frac{V}{U}\right]_{2} = \frac{1}{2} \left\{ 1 + \sqrt{1 - 4\frac{R_{2}}{R_{1}} \left[\frac{V}{U}\right]_{1} \left(1 - \left[\frac{V}{U}\right]_{1}\right)} \right\}$$
(3.1)

$$q_{j} = \left(A_{pos}I\right)^{2}R_{j} = I_{total}U\left(1 - \left[\frac{V}{U}\right]_{j}\right) = \frac{U^{2}\left(1 - \left\lfloor\frac{V}{U}\right\rfloor_{j}\right)}{R_{j}}$$
(3.2)

$$\frac{q_2}{q_1} = \frac{R_1 \left(1 - \left[\frac{V}{U}\right]_2\right)^2}{R_2 \left(1 - \left[\frac{V}{U}\right]_1\right)^2}$$
(3.3)



Figure 3.3 Change in heat rejection requirement from increases in resistance for batteries with different designed voltages at rated power.

The ratio of resistances may be found by equating the power for the two cases. Then the resistances, and areas if the ASIs are equivalent, are determined solely by the fraction of the open-circuit voltage at which they achieve rated power, Eq 3.4. Then substitution will give the

ratio of heat production at rated power for the two cases, Eq. 3.5. A battery that achieves rated power at 80 % of OCV will have a heat production at rated power that is 2.3 times higher than one designed at [V/U] = 90 %. A battery producing power at 70 % of the OCV will have 3.9 time higher heat generation than at [V/U] = 90 %.

$$\frac{R_{1}}{R_{2}} = \frac{ASI_{powerl}A_{pos2}}{ASI_{power2}A_{posl}} = \frac{\left[\frac{V}{U}\right]_{1}\left(1 - \left[\frac{V}{U}\right]_{1}\right)}{\left[\frac{V}{U}\right]_{2}\left(1 - \left[\frac{V}{U}\right]_{2}\right)}$$

$$\frac{q_{2}}{q_{1}} = \frac{\left[\frac{V}{U}\right]_{1}\left(1 - \left[\frac{V}{U}\right]_{2}\right)}{\left[\frac{V}{U}\right]_{2}\left(1 - \left[\frac{V}{U}\right]_{2}\right)}$$
(3.4)
$$(3.5)$$

Two different design changes would enable operating a battery at 90 % of OCV compared to 80 % while maintaining the same power output. First, a second identical battery may be connected in parallel to the original battery. This will lower the resistance of the battery pack by one half but will also double the energy and cost of the battery. A more cost-effective approach is to reduce the electrode thickness by coating a larger separator area with the same amount of active material. The capacity of the cell is maintained while minimizing increases in cost from a larger separator, current collector and packaging area. This approach is feasible as long as the reduced electrode thickness is above that at which the ASI begins to increase, which is around 20 microns for typical Li-ion electrode systems as discussed in detail below.

The efficiency of a battery defines the heat rejection requirements and may be measured or calculated. Measurement of round-trip efficiency of a battery is best performed by using a calorimeter to measure the heat given off during the cycling of the battery. The calorimeter removes the requirement of knowing the exact SOC of a battery during the entire drive cycle. Calculation of the round-trip efficiency of a battery requires a detailed transient battery model within a vehicle simulation program to exercise the battery over the many acceleration and deceleration periods that occur during a drive cycle. The interesting result is that the same battery will have different power ratings depending on what level of round-trip efficiency the user is willing to accept.

Figure 3.4 shows the efficiency of a battery as a function of the designed potential at which the battery reaches rated power. The figure is created using Equations 3.1 and 3.4 above. Each line may be considered a different drive cycle, or duty load, for a battery with the same energy but different impedance (changing separator area). The straight, solid black line represents the efficiency of the battery operated only at rated power, $P/P_{max} = 1$. In example, a battery designed at [V/U] = 0.8 will have 80 % efficiency for a single discharge pulse at rated power. Likewise, a battery designed at 0.9 will be 90 % efficient at rated power. Batteries are normally operated in the area above the line of the rated power. Therefore, the other curves represent the efficiency of

discharging a battery at power levels below rated power (typical driving conditions). Consider two batteries each designed for a rated power of 100 kW although one achieves this power at a [V/U] = 0.9 and the other at 0.7. If the two batteries are discharged at 45 kW, $P/P_{max} = 0.45$, the battery designed at [V/U] = 0.9 will be 6.4 % more efficient. This is significantly less than the 20 % efficiency improvement realized when operated at rated power. The efficiency penalty is reduced as the battery operates less and less near the rated power.



Figure 3.4 Efficiencies for batteries designed to achieve rated power at different fractions of their open-circuit voltage. Comparative efficiency lines are shown for equivalent power demands over a period of battery operation.

3.3 Governing Equations

The five coupled, algebraic equations that govern the battery design are presented in this section. While these equations are perhaps the most important, many other equations are used to fully define the battery mass and volume. These other equations will be specified where necessary in the following subsections.

The user of the model specifies the required maximum rated power, P_{batt} , of the battery. This power is translated to a current density, *I*, in Eq 3.6 using the area of the positive electrode, A_{pos} , the number of cells, N_{cell} , the open-circuit voltage at the SOC for power, $U_{ocv,P}$, and the fraction of the open-circuit voltage at which the designed power is achieved, [V/U].

$$I = \frac{P_{batt}}{A_{pos} N_{cell} U_{ocv,P} \left[\frac{V}{U}\right]}$$
(3.6)

The relationship between capacity and battery energy is described by Equation 3.7. Formally, the energy of a battery is the product of the capacity and the average voltage at which the energy is obtained. The average cell voltage is approximated in Eq. 3.7 by subtracting the polarization from discharging the battery at a C/3 rate from the open-circuit voltage at the SOC for energy, $U_{ocv,E}$. The energy for all batteries designed by the model is calculated at a C/3 rate and the average open-circuit voltage at 50 % SOC. The remaining necessary values are the capacity of the cell, *C*, ASI for energy, *ASI*_{energy}, number of cells, and area of positive electrode. Either the battery energy or capacity may be specified. The energy usage rate for the vehicle (Wh/mile).

$$E = N_{cell} C \left(U_{ocv,E} - \frac{C}{3} \frac{ASI_{energy}}{A_{pos}} \right)$$
(3.7)

The area of the positive electrode in Eq. 3.8 is determined largely by the area specific impedance for power, ASI_{power} , and resulting voltage drop. The voltage of cell at rated power, $V_{cell,P}$, is found from the product $[V/U]U_{ocv,P}$, where $U_{ocv,P}$ is the open circuit voltage at the SOC for max power. In general, the area of the electrodes will increase if the ASI for power increases. The areas of the negative electrode and separator are determined from the area of the positive electrode. The negative electrode is taken to be 1 mm larger than the positive electrode in both height and width to alleviate concerns of lithium plating during charge pulses. The separator area is slightly larger than the negative electrode to prevent the electrical shorting of the two electrodes.

$$A_{pos} = \frac{ASI_{power}P_{batt}}{N_{cell} (U_{ocv,P})^2 \left[\frac{V}{U}\right] \left(1 - \left[\frac{V}{U}\right]\right)}$$
(3.8)

The positive electrode thickness, L_{pos} , in Eq. 3.9 is determined from the capacity of the cell, *C*, specific capacity of the electrode material, *Q*, volume fraction of active material, ε_{act} , bulk

density of the active material, ρ , and the positive electrode area. The negative electrode thickness is determined by its specific reversible-capacity and the designed excess-capacity to prevent lithium plating during charging. We have chosen a ratio of 1.25 negative to positive reversiblecapacity (N/P ratio) for the default value for the cells with graphite negative electrodes. LTO negative electrode based cells are designed at a 1.1 N/P ratio because of the previously mentioned minimal possibility of lithium deposition. The maximum allowable electrode thickness is a user defined value. The calculation for the electrode area changes when the designed thickness is greater than the maximum allowed (Section 3.6.1).

$$L_{pos} = \frac{C}{Q\rho\varepsilon_{act}A_{pos}}$$
(3.9)

Finally, the ASI for power (and for energy to a lesser extent) is calculated using an expression that is based on the electrode thicknesses, the current density, and the C-rate. The exact expression will be discussed in the next session. The ASI in Eq 3.10 shows the basic dependencies with α and β being constant valued parameters.

$$ASI = \frac{\alpha + f(I)}{L_{pos}} + \beta$$
(3.10)

3.4 Calculation of the ASI

In most battery design scenarios, the ASI and [V/U] directly determine the electrode thickness and area to meet a specified power-to-energy (P/E) ratio and capacity requirement. Clearly, the ASI plays a significant role in the design of a battery and particularly in the case of the P/E ratios required by automotive applications. However, the ASI is not an inherent constant of a specific battery chemistry or cell design. The measured value of the ASI is a complex combination of resistances within the battery resulting from the physical processes occurring at different length and time scales. Consequently, the measured value is a function of many factors (state of charge, pulse length, current density, C-rate, particle size, transport and kinetic parameters, etc). The calculation used for the ASI in this battery design model has been discussed in detail and validated against experiments elsewhere.²⁰ The physical meaning of the equation will be discussed but those interested in the derivation are directed to the separate publication. We note that the ASI described here is slightly different than the one addressed in the paper. The thermodynamic component is removed that originated from the change in open-circuit potential with concentration for the intercalation materials. Equation 3.11 contains the definition of the ASI used in this document. I_{t1} is a positive valued current density for a discharge pulse. I_{t2} is equal to zero as it is during the relaxation period after the pulse. The subscripts are as follows: time 0, t0, is the time just before a current pulse begins, time 1, t1, is the time just before the current pulse ends, and time 2, t2, is the time long after the current pulse when the cell is at opencircuit and the concentration gradients have relaxed. Therefore, this ASI measurement is not troubled by accounting for a change in open-circuit voltage with the passage of current. In general, the ASI measured with this definition is similar, although smaller, in value to those produced using the more standard definition used elsewhere.

$$ASI = \frac{V_{t2} - V_{t1}}{I_{t1} - I_{t2}}$$
(3.11)

The ASI for the electrochemical charge and discharge process is referred here-in as ASI_{echem} . Our calculation approach for both the ASI for power and for energy involves adding three components together to reach the ASI_{echem} , Eq. 3.12. The first two factors include impedance that arises from the interfacial charge transfer and transport. The third factor is a lumped parameter used to capture the remaining impedance.

$$ASI_{echem} = ASI_{intf}^{pos} + ASI_{intf}^{neg} + ASI_{const}$$
(3.12)

The interfacial impedance for positive and negative electrodes both contain the charge transfer resistance component $\overline{R} T/(i_o a LF)$ as shown in Eq. 3.13 and 3.14. Here, i_o is the exchange current density related to the interfacial area and a is the ratio of interfacial area to electrode volume. An approximation often used for a relates the parameter to the volume fraction of the active material and the particle radius, $a = 3\varepsilon_{act}/r_p$. The variables i_o and a should be specified to relate to the same area as they are often not independently determined. \overline{R} and T correspond to the universal gas constant and absolute temperature respectively. F is Faraday's constant. The influence of the interfacial impedance is that the ASI_{echem} increases as the electrode thickness is reduced. This behavior is typically observed at electrode thicknesses less than 30 microns for common Li-ion battery materials.

$$ASI_{\text{intf}}^{\text{neg}} = \frac{\overline{R}T}{i_o a L_{neg} F}$$
(3.13)

$$ASI_{intf}^{pos} = \frac{\overline{R}T}{i_o aL_{pos}F} \left\{ \left(1 - \frac{I}{I_{lim}^{ionic}}\right) \left[1 - \left(\frac{r_c}{r_{c,lim}}\right)^2\right] \right\}^{-0.5}$$
(3.14)

The positive electrode interfacial impedance also includes two factors that account for the physical limitations that occur from depleting the concentration of the reactants within the porous electrode. The I_{lim}^{ionic} term is the limiting ionic current for lithium cation transport through the porous separator. The $r_{C,\text{lim}}$ term is the limiting C-rate for solid state diffusion of lithium in the active materials. The C-rate may be related to the current density with Eq. 3.15. Here, the specific capacity, Q, the active material density, ρ , active material volume fraction, ε_{act} , and the electrode thickness, L, are used. If either the limiting C-rate or limiting ionic current are approached, the ASI will begin to approach an infinite value. This approach assumes the cell and material design is such that the transport limitations all occur on the positive electrode. The parameters required for the ASI expression are fit to experimental measurements. The ASI values are corrected for the interfacial contributions present during measurement so that the correct ASI may be determined at different electrode thicknesses. This is termed "ASI correction factor" on the System Selection worksheet of the spreadsheet model.

$$I = r_C Q \rho \varepsilon_{act} L_{pos} \tag{3.15}$$

The cell ASI for energy, ASI_{energy} , and power, ASI_{power} , are determined by adding the ASI_{echem} to that of the current collectors, ASI_{cc} , as discussed in the next subsection. The difference between ASI_{energy} and ASI_{power} is that the limiting currents are not important during the C/3 discharge for energy and the ASI_{const} is a different value for two cases. ASI_{energy} will always be higher than ASI_{power} if a battery is operated far from the limiting current. The higher impedance is due to the formation of significant concentration polarizations during the longer time scale of the energy discharge. A reasonable rule-of-thumb is that the ASI_{const} for energy is 2.2 times the value for power in layered oxide materials such as $LiNi_{0.80}Co_{0.15}Al_{0.05}O_2$.

3.4.1 Current Collection Resistance

The resistance from the conductors used to collect the current must be accounted for as they can contribute significant ohmic drop to the battery. The ASI used to calculate the required cell separator area, ASI_{power} , is larger than the ASI for the electrochemical charge and discharge processes, $ASI_{echem,P}$, as shown in Equation 3.16. The ASI_{echem} value is typically measured from experiments and must be added to the external resistances that arise from the materials used to conduct the electric current. These resistances come from current collection in the cell and also those on the module and battery pack level.

$$ASI_{power} = ASI_{echem,P} + ASI_{cc} + ASI_{term}^{cell} + \frac{R_{cnct}A_{pos}}{N_{cells}}$$
(3.16)

The current collector foil impedance, ASI_{cc} , is determined from an analytical expression, Eq. 3.17, which accounts for the coated and uncoated region of the foil, labeled *act* for active and *tab* respectively. The resistance factor, R_f , and the resistance of the current collector foils, R_{cc} , are also shown for clarity in Eq 3.18 and 3.19. The factor of 2 in the R_f term is due to assuming half of the foil thickness carries the current produced on one side of the foil. While all of the current passes through the tab region, the magnitude of the current varies along the height of the coated foil as the reaction area continually contributes current to the foil. An equivalent length for the resistance calculation may be determined so that multiplication by the total current for a cell will give the correct ohmic drop. This equivalent length is H/3 if the current density is relatively constant over the entire area. The derivation of this equivalent length as well as an in-depth discussion of the voltage and current distribution in the foils may be found in subsection 3.4.2. Also in the later subsection, the assumption of constant current density is verified with numerical modeling.

$$ASI_{cc} = H_{act}W_{act}R_{cc} = R_f \left(\frac{(H_{act})^2}{3} + H_{act}H_{tab}\right)$$
(3.17)

$$R_{f} = \left(\frac{2}{\sigma_{foil,neg}L_{foil,neg}} + \frac{2}{\sigma_{foil,pos}L_{foil,pos}}\right)$$
(3.18)

$$R_{cc} = R_f \left(\frac{H_{act}}{3W_{act}} + \frac{H_{tab}}{W} \right)$$
(3.19)

The cell terminals are ultrasonically welded to the ends of the current collector foil tabs. While the welding removes this contact resistance, the ASI of the terminal must be included in the total cell resistance. The ASI of the cell terminals, ASI_{term}^{cell} , is the summation of the positive and negative cell terminals as shown in Eq 3.20. The dimensions for these terminals are set by the calculated width of the cell and the user defined terminal thickness and height.

$$ASI_{\text{term}}^{\text{cell}} = \left(\frac{1}{\sigma_{term,neg}} + \frac{1}{\sigma_{term,pos}}\right) \frac{H_{term}}{W_{term}L_{term}} A_{pos}$$
(3.20)

The ASI for connection losses is the last term in the ASI summation stated in Eq. 3.16. This ASI value is calculated by multiplying the ratio of cell positive electrode area to number of cells by the summation of the resistances, R_{cnct} , for cell terminals, module terminals, module interconnects, and batteries terminals. In this way, each cell shares in the burden of overcoming the system losses from carrying the electric current. The calculation of R_{cnct} is detailed in Eq. 3.21 with the individual sources of connection losses shown. The voltage drop resulting from cell-to-cell contact resistance, R_{cnct}^{cell} , is taken to be $10^{-4}U_{ocv,E}$ in Eq. 3.22, a small fraction of the open-circuit voltage. A battery manufacturer would only tolerate a minimal voltage drop from cell-to-cell contact. One connection method is to physically press the two cell terminals together. This resistance could be lowered by increasing the physical pressure and contact area, or by laser welding the terminals together. Regardless, the value used in the model is left to the choice of the user to leave as is or to change to a different value.

$$R_{cnct} = N_{cell} R_{cntct}^{cell} + N_{mod} R_{term}^{mod} + (N_{mod} - 1) R_{incnt}^{mod} + R_{term}^{batt}$$
(3.21)

$$R_{cnct}^{cell} = 10^{-4} \frac{N_{cell} U_{ocv,E}}{IA_{pos}}$$
(3.22)

The module terminal resistance, R_{term}^{mod} , calculation in Eq. 3.23 is shown as an example of how the terminal and interconnect resistances are calculated for the module and battery pack. The size of the terminals and thus their resistance are determined from a calculation based on a predetermined allowable rate of temperature rise for the conductor. This approach is explained in more detail in subsection 3.4.3.

$$R_{\text{term}}^{\text{mod}} = \frac{H_{term}}{\sigma_{term} \pi (r_{term})^2} N_{term}^{\text{mod}}$$
(3.23)

3.4.2 Potential and Current Distribution in the Current Collection Foils

The designed current collection system was evaluated using a numerical simulation package. Equations 3.24-3.26 were solved for a steady state, isothermal, and 1-D simulation. Here, the conductivity, σ_j , is the effective conductivity of $\frac{1}{2}$ of the foil (the other half carries the current from the opposite side). The bulk conductivity value, σ_j^0 , is multiplied by the thickness of the conductor, $L_j/2$, to lower the dimension of transport.

$$I_n = \frac{U_{ocv} - \left(\Phi_{1,pos}(x) - \Phi_{1,neg}(x)\right)}{ASI_{echem}}$$
(3.24)

$$\nabla \cdot \left(\sigma_{pos} \nabla \Phi_{1,pos} \right) = -I_n \tag{3.25}$$

$$\nabla \cdot \left(\sigma_{neg} \nabla \Phi_{1,neg}\right) = I_n \tag{3.26}$$

The boundary conditions were set for both ends of each foil. The tab ends of the foils were set to a specified voltage and the opposite ends of the foils were restricted to a no flux condition. The simulation was performed using the foils defined in our battery design: 12 micron thick copper foil and 20 micron thick aluminum. The cell length was 20 cm, the ASI_{echem} was 30 ohm cm², and the U_{ocv} and V_{cell} were set to 3.72 and 3.57 V respectively. Figure 3.5 shows the current and potential distribution in the foils and in the cell resulting from the simulation. The cell potential along the length of the foil varies only by 1.5 mV from maximum to minimum difference. The 0.04 % variation in voltage results in a 0.9 % variation in current density. This verifies the current density is uniform along the length of the foil. This is also obvious from the linear relationship of current with foil height in Fig. 3.5. The assumption of constant current density was tested in cell heights up to 100 cm and found to be satisfactory. The assumption should be reasonable as long as the ASI_{echem} is at least twice the value of ASI_{cc} . The simulated resistance of the foils is found to raise the ASI_{echem} by 0.7 for an ASI_{power} of 30.7 ohm cm². Additionally, the numerical result verified that H/3 is the correct equivalent length to represent the ASI_{cc} for the cell. This may also be found analytically, Eq. 3.27-3.29, if you assume an even current distribution. We have shown that to be a reasonable assumption.

$$\Phi_{1,pos} = V_{cell} + \frac{I_n}{2\sigma_{pos}} \left(H^2 - x^2 \right)$$
(3.27)

$$\Phi_{1,neg} = \frac{I_n}{\sigma_{neg}} \left(\frac{x^2}{2} - xL \right)$$
(3.28)

$$ASI_{echem} + ASI_{cc} = \frac{1}{H} \int_{0}^{H} \frac{\Phi_{1,pos} - \Phi_{1,neg}}{I_n} dx = \frac{V_{cell}}{I} + \frac{H^2}{3} \left(\frac{1}{\sigma_{pos}} + \frac{1}{\sigma_{neg}} \right)$$
(3.29)



Figure 3.5 The change in current and potential within the positive and negative foils. The current collection design results in a uniform current distribution along the length of the foil.

An analogous problem has been solved by Euler and Nonnemacher and then communicated repeatedly by Newman *et al.*^{24, 25} The analytical solution they presented may be used after a slight alteration to dimensionalize the current density to the geometry of our concern, Eq. 3.30 and 3.31. This solution was reached assuming linear polarization behavior and is valid for cases where the current density varies along the height of the current collector foil. Thus, this approach is a more general solution than the one we use in the design model.

$$ASI_{echem} + ASI_{cc} = \frac{U_{ocv} - (\Phi_{1,pos} - \Phi_{1,neg})}{I} = \frac{H^2}{\sigma_{neg} + \sigma_{pos}} \left[1 + \frac{2 + \left(\frac{\sigma_{neg}}{\sigma_{pos}} + \frac{\sigma_{pos}}{\sigma_{neg}}\right) \cosh v}{v \sinh v} \right] (3.30)$$

$$v^{2} = \frac{H^{2}}{ASI_{echem}} \left(\frac{1}{\sigma_{pos}} + \frac{1}{\sigma_{neg}} \right)$$
(3.31)

3.4.3 Determination of Module Terminal, Battery Terminal, and Module Inter-connect Size

An important factor for setting the resistances of a module terminal, battery terminal, or module interconnect is the allowable rate of temperature rise in the conductor at full power. We set the acceptable rate of temperature rise, dT/dt, at 0.2 °C/sec or a 2 °C rise for a 10-sec power burst under adiabatic conditions. The heating rate, q, is then used to determine the mass, m, of the terminal required for the designed battery in Eq. 3.32. Since the heating rate may also be determined by Eq. 3.33, we may determine the cylindrical terminal radius and mass by assuming a length, H_{term} . In this way, the size of the module terminal is redesigned during each simulation to meet the specified power requirements and allowable temperature rise, Eq. 3.34. The mass of the conductor is found to be inversely proportional to the allowable temperature rise.

$$q = mC_p \frac{dT}{dt} \tag{3.32}$$

$$q = \frac{H_{term}}{A_{term}\sigma_{term}} (IA_{pos})^2$$
(3.33)

$$A_{term} = IA_{pos} \left[\rho_{term} \sigma_{term} C_p \frac{dT}{dt} \right]^{-1/2}$$
(3.34)

A copper busbar must also be sized for batteries using a single row of modules. We have somewhat arbitrarily assumed a $\Delta V_{bb} = 30$ mV drop across the busbar to be allowable at maximum current. This value maybe easily changed by the user. Equation 3.35 is used to calculate the mass of the busbar, m_{bb} . The complicated expression for the volume of the busbar is derived from the voltage drop, conductivity, busbar width, w_{bb} , and required busbar cross-sectional area.

$$m_{bb} = \rho_{bb} \frac{I_{tot} (w_{bb})^2}{\Delta V_{bb} \kappa_{bb}}$$
(3.35)

3.5 Calculation of Battery Dimensions

The goal of the model is to quantify how the various components of a specific battery design sum to make the mass and volume of the battery pack. In this way, a true energy and power density can be calculated as well as the exact materials requirement to meet this design. Summing the mass of the components is relatively straight forward. Determining the total volume that contains the components and required free volume is not as obvious. The exact calculations used in the design model are detailed below for the cell, module, and battery pack.

3.5.1 Cell Dimensions

The number of layers in each cell is approximated in Eq. 3.36 by accounting for the compression factor, X_{comp} , and the individual thicknesses of the current collector foils, L_{foil} , electrodes, L_{pos} and L_{neg} , separator, L_{sep} , and container, L_{cont} . X_{comp} is usually taken to be 0.97. The Li-ion battery chemistries this model was designed for are assumed to undergo negligible volume change on the cell level. Slight volume changes can be accommodated as mentioned in Section 2.3.

$$N_{layers} = X_{comp} \frac{L_{cell} - 2L_{cont} + L_{foil}^{neg}}{L_{foil}^{neg} + L_{foil}^{pos} + 2(L_{sep} + L_{neg} + L_{pos})}$$
(3.36)

The N_{layers} approximation is necessary as the cell thickness is a user defined parameter. The aspect ratio of the cell is also user defined; therefore, solving for the width also determines the height of the cell as seen in Eq. 3.37. The width is calculated from the number of layers and the aspect ratio, H/W. The factor of 2 enters the denominator as both sides of the foil are assumed to be coated.

$$W_{pos} = \sqrt{\frac{A_{pos}}{2\frac{H}{W}N_{layers}}}$$
(3.37)

Having determined the width and height of the electrode, the rest of the cell dimensions are relatively straightforward, Eq. 3.38 and 3.39. The width of the cell, W_{cell} , is 2 mm wider than the positive electrode to allow for the larger separator area and pouch seals. The thickness of the folded edge, L_e , is also included in the width dimension. The pouch seals are folded up, pressing along the inside wall of the module casing. The height of the cell, H_{cell} , is the height of the positive electrode in addition to the distance for the terminals and connections to the foil tab, $L_{term,cnt}$. Our assumed design requires 15 mm for this distance at each end of the cell for a total of 30 mm. The volume of the cell is the product of the three dimensions.

$$W_{cell} = W_{pos} + 2 + 2L_e \tag{3.38}$$

$$H_{cell} = H_{pos} + 2L_{term,cnt} \tag{3.39}$$

3.5.2 Module Dimensions

The module dimensions are defined by Eq. 3.40-3.42. The height and length of the module are both just 2 mm wider than the cell dimension. The width of the module is related to the total thickness from all of the cells with allowance for a SOC controller at one end.

$$H_{\rm mod} = W_{cell} + 2 \tag{3.40}$$

$$L_{\rm mod} = H_{cell} + 2 \tag{3.41}$$

$$W_{\text{mod}} = L_{cell} \left(N_{cell/\text{mod}} + 1 \right) + 1 \tag{3.42}$$

3.5.3 Battery Pack Dimensions

The battery pack volume includes all of the modules, spacing for connections between modules, channel for the cooling air to flow, H_{air} , thickness of the module compression plates, L_{comp} , and the battery pack jacket, L_{jack} (Eq. 3.43-3.45). L_{jack} includes a 10 mm thick insulation layer sandwiched between two aluminum walls for the container. The thickness of the aluminum wall increases from 1 to 1.5 to 2 mm as the battery volume increases from < 20 L to < 40 L to larger dimensions. The layout of the modules, number per row, $N_{mod/row}$, and number of rows, N_{row} , is also included. The final volume of the battery is the product of the three dimensions. The space left for connections between modules, L_{gap} , is a function of the number of rows of modules. L_{gap} is equal to 8, 10, or 20 depending if there is one, two, or four rows of modules. Three rows of modules are not allowed as the positive and negative terminal for the battery would be on opposite ends and thus not very practical. A number greater than four rows of modules is deemed unnecessary.

$$H_{batt} = H_{mod} + 2H_{air} + 2L_{jack}$$
(3.43)

$$L_{batt} = N_{\text{mod/row}} W_{\text{mod}} + H_{air} + 2L_{comp} + 2L_{jack}$$
(3.44)

$$W_{batt} = N_{row} L_{mod} + L_{gap} + 2L_{jack}$$
(3.45)

3.6 Maximum Electrode Thickness

A practical limitation exists for the maximum achievable electrode thickness. This limitation may be set by manufacturing capabilities, ionic and electronic current transport within the porous electrode, susceptibility to plating lithium on the negative electrode, or aging characteristics related to adhesion to the current collector. Some of these challenges are discussed in great detail in the following subsection. When the maximum electrode thickness, L_{max} , has been reached on either the positive or negative electrode, the electrode area equation is modified as shown in Eq 3.46. The electrode thickness, L_{tgt} , is the largest electrode thickness, negative or positive, calculated at the targeted fraction of the OCV [V/U].

$$A_{pos}^{L\max} = \frac{L_{tgt}}{L_{\max}} A_{pos}$$
(3.46)

The area of the electrode is now determined by the cell capacity requirement to meet the battery energy demands and not the target voltage at rated power. As a consequence, the battery pack will operate at a higher [V/U] than originally selected by the battery designer. The new [V/U] may then be calculated from Eq. 3.47 which is the solution to the quadratic found in Eq. 3.48.

$$\left[\frac{V}{U}\right] = \frac{1}{2U_{cell}} \left[U_{cell} + \sqrt{\left(U_{cell}\right)^2 - 4\frac{P_{batt}ASI_{power}}{N_{cells}A_{pos}}}\right]$$
(3.47)

$$V_{cell} = U_{cell} - \frac{P_{batt}ASI_{power}}{V_{cell}N_{cells}A_{pos}}$$
(3.48)

The maximum electrode thickness may have a large impact on the energy density and cost of cells designed for high energy and range. Nelson et al. demonstrated this concept in 2009 assuming a 100 micron maximum electrode thickness.^{12,19} In 2010, Santini et al. relaxed this assumption to 300 microns; although, the thickest electrode discussed in the paper was a 225 micron graphite electrode in the LMO-Gr EV with 100 mile range.¹³ In conversations with manufactures, 100 microns appears to be the general electrode thickness used for EV type cells at the present time. However, Santini et al. has shown substantial increases in energy density and decreases in cost if larger electrode thicknesses may be utilized. The challenges to achieving thick electrodes, in addition to those already mentioned, relate to fast charging while avoiding lithium metal deposition, utilizing all of the materials reversible capacity, removing gases formed during formation cycling, wetting the full porosity of the electrode, achieving defect free coatings, and drying the thick electrode at high rates. Our opinion is that the successful cell manufacturers will engineer ways to overcome these challenges to increase energy density and lower cost.

Dependent upon the battery chemistry and designed P/E ratio, the maximum achievable electrode thickness (loading) may have a significant effect on the end cost and energy density of a battery pack. For batteries designed at low P/E ratios or for cell chemistries with low volumetric capacities, the designed electrode thickness based on the target efficiency is often larger than what is feasible during operation in a transportation environment. This subsection explores some of the challenges that arise in the electrochemistry when larger electrode thicknesses are utilized.

Argonne gained a wealth of experience in the NCA-Gr in $1.2 \text{ M LiPF}_6 3:7 \text{ EC:EMC}$ cell during the Advanced Technology Development program sponsored by the US Department of Energy (DOE). Dees and coworkers developed a world-leading parameter set for a numerical model

through exhaustive electrochemical measurements, ex-situ characterization techniques, and multi-scale modeling activities.²⁶⁻³⁵ The resulting phenomenological cell model founded on the methodology originating from John Newman (UC Berkeley) will be used to evaluate the electrochemical behavior of cells using thick electrodes.³⁶ The coupled, non-linear partial differential equations are solved with the finite element method using FlexPDE.

Simulated discharge capacity for the C/1 and C/3 discharge rate is shown in Figure 3.6 as a function of electrode thickness. For reference, the target positive electrode thicknesses for this cell operating at a 5C-rate and a [V/U] = 0.8 is 142 microns. The line of 100 % capacity utilization is also shown as a means to judge the deviation from theoretical capacity. As expected, the C/1 rate deviates more strongly than the C/3 rate with increasing electrode thickness. The loss in capacity is a result of the cell hitting the discharge voltage cut-off, 3.3 V, before all of the lithium has been transported from the negative to the positive electrode.

Figure 3.7 displays the normalized concentration profile of the electrolyte salt, LiPF₆, at the end of a C/1 and C/3 discharge for an electrode thickness of 245 microns. The C/1 discharge results in a positive electrode starved of electrolyte salt. This transport limitation results in the cell prematurely reaching the voltage cutoff. In order to overcome this limitation, the electrode would need to be engineered with significantly reduced tortuosity³⁷ or utilize an electrolyte with better mass transfer characteristics. This behavior is exacerbated by lower temperatures, such as those experienced during winter driving conditions. The fraction of theoretical discharge capacity begins to lower significantly at thicknesses greater than 100 microns, 3.4 mAh/cm², at the C/1 rate and 175 microns, 6.4 mAh/cm², at the C/3 rate. The electronic transport properties of the cathode material also play an important role in determining the current distribution within the electrode. While the NCA material has a reasonably high conductivity, other cathode materials have lower valued electronic conductivities and, depending on the conductive additive properties, may have different current distributions and limitations within the electrode.



Figure 3.6 Cell capacity simulated at the C/1 and C/3 rate as a function of electrode thickness (loading) for NCA-Gr.



Figure 3.7 Normalized electrolyte salt concentration at the end of discharge at the C/1 and C/3 discharge rates. The second half of the positive electrode next to the current collector is starved of the LiPF₆ salt resulting in a lower utilization of the inherent cell capacity.

The simulated ASI for a 5C, 10-s discharge pulse at 60 % SOC is shown in Figure 3.8 as a function of electrode thickness. The initial decrease in the ASI is a mathematical result of diminishing significance of the interfacial impedance as more current is passed in the same geometric area. The ASI then remains constant from 75 microns to nearly 400 microns. The constant ASI results from ohmic losses that behavior linearly with applied current. The dramatic increase in the ASI at the largest electrode thicknesses results from limitations in electrolyte transport within the porosity of the positive electrode. This is similar to what is displayed in Fig 3.7 above during the constant discharge at the C/1 rate for an electrode thickness of 245 microns.

The most significant issue for pulse power operation with thick electrodes occurs on the negative electrode during a charge or regen pulse, Figure 3.9. The potential of the negative electrode may drop below that of a hypothetical lithium reference electrode during a charge pulse, inferring an undesirable side reaction of lithium plating on graphite.³⁸ This behavior is exacerbated by increasing electrode thickness. Operation at higher SOC and lower temperatures will also increase the probability of lithium plating. The lithium reference electrode is taken to be in the center of the separator layer. The two times shown in the graph, 1-s and 10-s, represent different polarization measurements for the electrode. The 1-s value includes all of the interfacial impedance and minor contributions from concentration polarization. The longer time value includes additional changes in potential due to the concentration gradient in the electrolyte. The 1-s time is the more accurate valuation of the tendency of the electrode to plate lithium.



Figure 3.8 Calculated ASI from a simulated 10-s, 5C discharge pulse for the NCA-Gr cell couple at 60% SOC. Positive and negative electrode thicknesses are similar in value for this cell design. Transport within the electrolyte is not limiting until the electrode thickness approaches 450 microns for these simulation conditions.



Figure 3.9 The potential of the negative electrode versus a hypothetical lithium reference electrode located in the center of separator during a 5C charge pulse for the NCA-Gr couple.

The tendency for lithium plating appears to be significant at electrode loadings greater than 3.4 mAh/cm² under these specific conditions. Lower continuous and pulse charge rates are most likely necessary to ensure the safe operation and long life of this battery. Fast charging of PHEV and EV batteries is an oft discussed value-added characteristic necessary to increase the attractiveness of electric vehicles to the consumer. However, this fast charge requirement may require a lower loading design to prevent lithium plating from occurring. This would not be true for batteries based on LTO negative electrodes, or possibly even non-graphitic carbon electrodes (e.g. hard carbon). The consequence of a lower loading design is that higher quantities of inactive materials are used resulting in a more expensive and less energy dense battery.

A limit of 100 microns has been chosen for the default maximum electrode thickness. This thickness represents a graphite electrode balanced to a positive loading of 3.5 mAh/cm² and is the largest thickness that ASI measurements have been validated at Argonne. However, a low volumetric capacity electrode, such as LMO, will result in a lower area-specific capacity as the limit will be determined by the positive electrode thickness. One domestic OEM has suggested that at the time of this publication, state of the art electrode loadings for PHEV applications are less than 2 mAh/cm². This low loading level was selected based on cold-start performance, life testing, and rate capability studies. We note that different electrolytes will likely have the most significant effect on the transport limitations and result in different optimum electrode loadings. Gel-based electrolytes or standard carbonate electrolytes mixed with low molecular weight polymers are often utilized in pouch cell based batteries. While the ionic conductivity of these systems may approach standard carbonate electrolytes used, diffusion of the salt is restricted.

This decrease in mass transport will result in large increases in impedance during longer discharges (constant speed highway driving). Hence, companies using an electrolyte with sluggish mass transport will require thinner electrodes than what the model would normally calculate based on pulse power applications. Greater electrode thicknesses may be achievable in the future as manufacturers expend significant engineering efforts to minimize the inactive material that lowers energy density and raises cost. The concerns over lithium plating may remain unless new, less susceptible negative electrodes are developed that still enable high energy density.

Table 3.2 displays a calculated sensitivity analysis for designing the battery at different target electrode loadings to achieve 17 kWh of total energy for the NCA-Gr cell couple for two power levels, 120 kW and 60 kW of power. The consequence of varying electrode loading at constant power and energy is to change the fraction of open circuit voltage at which rated power is achieved at BOL. This quantitative comparison of electrode thickness with [V/U] is only valid for the specific battery and cell chemistry designed in the table, while the qualitative results would hold for all systems. An alternate but equivalent way to create Table 3.2 is to maintain a constant value of [V/U], but vary the designed power level. As mentioned previously, private communications suggest that current electrode thicknesses used in PHEV applications are near 50 microns or 2 mAh/cm²; however, the exact details of that cell configuration are unknown and thus direct comparison should be conducted with caution. For the cell chemistry shown in the table, a loading of 1.9 mAh/cm² with 120 kW of power correlates to achieving the rated power at BOL at [V/U] = 0.84. Our default suggestion of using 80% of OCV results in a moderately less expensive battery, albeit similar in value. If a battery of lower P/E ratio were designed, then the electrode thickness (loading) would be much larger for the same designed fraction of opencircuit voltage. The sensitivity to [V/U] decreases with lower [V/U] values owing to the inverse proportionality of electrode loading to the P/E ratio. The calculated designed electrode loadings for lower P/E ratios will differ the most significantly from those used in industrial practice today. This is in part the motivation to limit electrode thicknesses to 100 microns as well as the transport limitations discussed earlier. While the standard practices of today are important, the goal of the calculations is to evaluate the potential cost of Li-ion batteries in the future years after improvements have been made resulting from the competitive marketplace. Therefore, calculating a range of values will be the most instructive approach to determining where future battery costs may fall. Additional variation studies may be found in Appendix A.

60 kV				120 kW		
17 kWh	Loading	Thickness	Battery cost	Loading	Thickness	Battery cost
[V/U]	mAh/cm ²	μm	US\$	mAh/cm ²	μm	US\$
68	7.0	180	4218	3.2	82	4664
72	6.5	166	4244	3.0	76	4726
76	5.8	149	4284	2.7	68	4822
80	5.1	129	4345	2.3	58	4969
84	4.2	107	4445	1.9	48	5210
88	3.2	81	4647	1.4	35	5653
92	2.1	54	5039	0.9	22	6699

Table 3.2 The effect of electrode loading on the price of a 17 kWh NCA-Gr PHEV40 battery with 96 cells

4. Thermal Management

The power and life of Li-ion batteries are more drastically affected by temperature than most other batteries including those based on lead-acid, nickel-cadmium, and nickel metal-hydride systems. It is important that the temperature of a Li-ion battery be controlled at all times, even when the battery is at rest. Developing schemes for effectively controlling the pack temperature at minimum cost will certainly be important in the success of this technology. The most difficult problem is the removal of heat generated within the battery, principally by ohmic heating. Avoiding excessive temperature rise during idle periods in hot ambient conditions is also a problem. Either of these conditions might raise the temperature to well above 40°C, which enhances degradation reactions and shortens the battery life. In fact, maintaining the battery near the minimum temperature for adequate power will prolong battery life by retarding degradation reactions. Because the battery has poor power at low temperatures, heating the battery from a very cold condition is necessary and especially difficult for large EV battery packs for which no assistance is available from the engine. For electric-drive vehicles to be competitive in the market with conventional vehicles, these thermal control problems must be solved at moderate costs by means that do not compromise the safety of the vehicle or battery system.

The BatPaC model has a separate worksheet for designing the thermal management system. The results are transferred to the Battery Design worksheet to calculate the mass, volume and materials requirements for the battery pack.

4.1 Heat Generation Rates in the Battery Pack during Driving

During driving, the heat generation rate depends on the drive cycle and the power of the battery relative to the demands of the cycle for the vehicle being driven. As discussed below in section 4.2, fluctuations in the temperature of the pack are smoothed out by the battery heat capacity. The rate that the cooling system must handle is the average rate for the most difficult sustained driving conditions to which the battery pack will be subjected.

There are two types of driving profiles that will generate high heating rates in the battery pack. One is a drive cycle such as the US06 that has rapid acceleration and deceleration, including periods at high speeds. The average heat generation rate for such driving profiles is difficult to evaluate and is best done by vehicle simulation studies. These studies require a battery impedance algorithm that makes possible accurate estimates of internal battery heating during vigorous driving patterns. The results of vehicle simulation studies of battery heating can be entered on line 37 in the Thermal worksheet to override the estimated default values. A second driving profile that causes a high rate of battery heating is driving at a high, constant speed for a prolonged period (i.e. >5 minutes). Battery heating is easier to estimate for this condition because the battery power requirement is a direct function of the vehicle speed and the battery impedance approaches a constant value after a few minutes of driving.

Vehicle simulations were completed by Argonne's Advanced Powertrain Research Facility with the Autonomie model. The battery energy required to support all-electric driving was compared for the US06 driving cycle and a constant speed of 70 mph based on a vehicle similar to the Chevrolet Volt. In that study the impedance of the LMO-G battery was determined with a model

that had been derived from HPPC data. The model has an ohmic resistance and two polarization resistances with time constants of 15 and 270 seconds, respectively. The model is useful for calculating the changing impedance during a complex driving cycle such as the US06 cycle and it reproduces the impedance values entered in the "Chem" worksheet for the stated conditions. Table 4.1 shows the results of this study including the parameters derived to enable calculation of the heat generated in the battery operated at constant speed.

The first column of values shows the results of the study for the Autonomie model for both the US06 cycle and constant speed driving at 70 mph. The sum of the power for accessories, rolling friction and aerodynamic drag was divided by the total power determined in the Autonomie model to obtain a vehicle efficiency factor. This factor accounts for power losses in the power converter, the motor and the gears. This vehicle efficiency factor is assumed to be sufficiently constant over the vehicle speeds of interest for the purpose of establishing battery cooling requirements and the associated cost.

The energy demand per mile of travel for the vehicle of Table 4.1 was calculated to be 250 Wh/mile from the useable battery energy of 10 kWh for 40 miles of travel on the Urban Dynamometer Driving Schedule (UDDS). As shown at the bottom of the first column of figures, the estimated constant speed at which this energy demand would result is 57.85 mph. The second column of figures shows how the key parameters from the first column have been adjusted to round numbers that are suitable for the approximations required in BatPaC, and these values are highlighted in yellow. BatPaC then calculates the remaining parameters in close agreement to Autonomie.

The electric energy consumptions (Wh/mile) for the two types of driving are almost the same, but the higher average speed at 70 mph indicates a higher energy consumption per unit time (higher power) for this constant speed than for the US06 cycle (average speed of 48.4 mph). More importantly, a high heating rate results because constant rate discharge of the battery pack causes high battery impedance, which reaches more than twice the impedance of the battery when operated for short power bursts. The US06 cycle has long periods of high speeds resulting in elevated battery impedance, but this increase in impedance is mitigated by periods of low power demand during which the impedance decreases. Thus, driving at a constant high speed for an extended time results in the maximum heat generation rate in the battery and justifies our emphasis on this source of battery heating, which fortunately is the easiest to calculate.

For microHEV battery packs, heat generation is intermittent and substantial periods of little or no heat generation exist in the load profile. The model estimates the heat generation rate for 25-kW microHEV batteries at 100 W, which is entered on line 36 in the Thermal worksheet. For the larger battery packs (HEV-HP, PHEV and EV), the BatPaC model calculates the vehicle constant speed at which the centerline of the hottest cells in the battery will reach the maximum allowed temperature with steady state cooling, which is set at 35°C in the sample calculation on line 18 of the Thermal worksheet. That speed is deemed to be the maximum speed that the vehicle can be driven without damaging the battery by overheating. The battery control system should be designed to limit the battery output as this condition is approached.

Table 4.1 Energy Requirements for EREV Midsize 40AER similar to Chevrolet Volt. Highlighted values are the parameters required by BatPaC to estimate heat generation.

	Autonomie Values	Adjusted for BatPaC
Vehicle Parameters		
Mass, kg	1850	
Power for accessories, kW	0.472	<mark>0.5</mark>
Rolling resistance, kg/kg	0.008	
Factor for rolling resistance power, kW/mph	0.0647	<mark>0.065</mark>
Coefficient of drag, CD	0.311	
Frontal area, m ²	2.372	
Factor for aerodynamic drag, kW/(mph) ³	4.060E-05	<mark>4.0E-05</mark>
Parameters for LMO-G Battery		
Battery power, kW	100	100
Battery energy, kWh	15.0	15.0
Usable battery energy, kWh	10.0	10.0
Designated electric range, miles	40.0	40.0
Standard energy demand, Wh/mile	250	<mark>250</mark>
Autonomie Results		
US06 Cycle		
Electric energy consumption, Wh/mile	327.3	
Average power (excluding stopping and regeneration), kW	25.2	
Maximum power, kW	126.4	
70 mph Constant Speed		
Power, kW	22.67	
Electric energy consumption, Wh/mile	323.8	
Constant Speed Results from Vehicle Parameters		
70 mph		
Power for accessories, kW	0.472	0.5
Power for rolling friction, kW	4.53	4.55
Power for aerodynamic drag, kW	13.93	13.72
Vehicle efficiency factor*	0.835	<mark>0.833</mark>
Total power, kW	22.67	22.53
Electric energy consumption, Wh/mile	323.8	321.9
Speed for 250 Wh/mile Energy Demand		
Estimated Speed (S_o), mph	57.85	<mark>58.0</mark>
Power for accessories (P_a) , kW	0.472	0.5
Power for rolling friction (P_f), kW	3.74	3.77
Power for aerodynamic drag (P_d) , kW	7.86	7.80
Vehicle efficiency factor $(e)^*$	0.835	0.833
Total power (P_o) , kW	14.46	14.49
Electric energy consumption, Wh/mile	250	250

*Determined by dividing the sum of the power for accessories, rolling friction and aerodynamic drag by the total power determined in the Autonomie model.

This maximum speed will depend on vehicle design parameters, the type of electric drive (HEV-HP, PHEV or EV), the battery maximum power, the type of cooling system, (cabin air or liquid) and the battery design as it affects the efficiency of cooling. Under some conditions, it might be desirable to set a lower speed than that limited by battery heating to reduce the required capability and cost of the cooling system. This can be done by entering the maximum constant speed at which the vehicle will be operated on line 20 and BatPaC will calculate the maximum steady-state temperature at the center of the cells on line 19.

Since it has been established that the maximum battery heating will occur when the vehicle is driven at a high, constant speed, it is necessary to estimate the power needed to drive the vehicle at any desired speed and the resultant battery heating. For this purpose, approximate equations are needed for vehicle parameters including accessory power, rolling friction and aerodynamic drag. A starting point for deriving these equations is the battery power calculated from the energy requirement per mile, \dot{E} (Wh/mile) entered on the Battery Design worksheet, line 172. The battery power, P_{s0} (kW), will sustain a vehicle speed of S_0 at the designated value of \dot{E} .

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$$P_{s0} = S_0 \cdot E \ /1000 \tag{4.1}$$

The total battery power for maintaining the designated speed for the energy requirement is the summation of power demands for accessory power (P_a), rolling friction power (P_f), aerodynamic drag power (P_d) and power losses resulting from inefficiencies in the electrical and mechanical components of the powertrain system. From the calculations described in Table 4.1 we estimate the efficiency (e) from the battery to the wheels and accessories to be approximately constant at 0.833. The accessory power is estimated to be constant at 0.5 kW, but the model user may wish to increase this value to allow for air conditioning in warm climates and battery cooling. Estimates of the power to the system for cooling the battery (based on a coefficient of performance of 2.5 for the refrigeration cycle) are calculated on line 39 of the Thermal worksheet.

For midsized sedans, the power to drive the vehicle can be estimated with the energy requirement per mile, (Wh/mile) as the starting point. A reduction in the future from the default value of 250 Wh/mile will result primarily from reduction in the vehicle weight, which will reduce rolling friction. Thus, for the rolling friction factor, f_{Pf} (kW/mph), we propose:

$$f_{Pf} = 0.065 \cdot \dot{E} / 250 \tag{4.2}$$

And, the power for rolling friction, $P_f(kW)$ is:

$$P_f = f_{Pf} \cdot S \tag{4.3}$$

Aerodynamic drag is more difficult to reduce than vehicle weight. However, the interaction between the speed and the total power (a function of both P_f and P_d) affects the drag factor, f_{Pd} , which is the product of the drag coefficient and the vehicle frontal area. The drag factor is estimated here from the default value of 0.00004 kW/(mph)³.

$$f_{Pd} = 0.00004 \cdot (\dot{E}/250)^{0.3} \tag{4.4}$$

The value of P_d at any vehicle speed, S, may be calculated with the expression:

$$P_d = f_{Pd} \cdot (S)^3 \tag{4.5}$$

The battery power, P_s , at any specified sustained vehicle speed can then be calculated from:

$$P_{s} = (P_{a} + P_{f} + P_{d})/e$$
(4.6)

Eq. (4.1) and (4.6) are solved simultaneously for S_0 by iteration on lines 8 and 9 of the Thermal worksheet. S_0 is used to calculate f_{Pf} and f_{Pd} by means of Eq. (4.2) and (4.4).

For battery packs other than microHEV battery packs, the BatPaC model calculates the heat generation rate, q (W), from the battery current, I_{total} (A), and the battery resistance for a sustained constant discharge, R_{bs} (ohms).

$$q = (I_{total})^2 \cdot R_{bs} \tag{4.7}$$

The value of R_{bs} for PHEV and EV batteries is calculated from the total ASI for the energy of the cell (taken at the C/3 rate and an average resistance assumed to be the same as that at 50% SOC), and entered on line 28 of the Thermal worksheet. This value is equivalent to the sum of the ohmic and polarization resistances for the impedance model used in the Autonomie vehicle simulation study described above. For HEV-HP applications, the battery is operated for an estimated 30 to 60 seconds and the value of R_{bs} is estimated to be 20% more than for the battery resistance at maximum power. Also the average battery heat generation rate is conservatively estimated as 60% of q as defined by Eq. (4.7) because only about 60% of the power for the vehicle is delivered from the battery; the balance of the power for the vehicle comes from the engine by direct coupling to the drivetrain or through a generator and motor. For PHEV and EV batteries, the heat generated in the battery pack is calculated directly from Eq. (4.7). Battery current and voltage, V, are calculated as follows:

$$I_{total} = 1000 \cdot P_s / V \tag{4.8}$$

$$V = \frac{-U_{ocv,E} + \sqrt{(U_{ocv,E})^2 - 4 * 1000 * P_s * R_{bs}}}{2}$$
(4.9)

The heat generation rate in the battery is selected on line 38 in the Thermal worksheet from three possible values: (1) for microHEV batteries it is 100 W (line 36), (2) for batteries that have been modeled in a vehicle simulation program to determine maximum sustained heat generation and the result entered on line 37, and (3) for all other battery packs, the heat generation rate calculated on line 34 (based upon Eq. 4.7 and modified as discussed above for HEV-HP batteries) is selected.

As a check on the practicality of estimating the heat generation in the battery by the method described above, another calculation was done by Argonne's Advanced Powertrain Research

Facility with the Autonomie model. In that calculation, the vehicle was an advanced sedan with an energy demand of only 201.8 Wh/mile on the UDDS cycle resulting from a vehicle weight of only 1625 kg, rather than 1850 kg as in the initial study (Table 4.1). The drag factor was only slightly lower than previously (i.e. 3.7×10^{-5} vs. 4.0×10^{-5}). A comparison of the results of calculations on Autonomie and BatPaC are shown in Table 4.2.

	Autonomie	BatPaC
Energy demand (\dot{E}),Wh/mile	201.8	201.8
Constant speed at \dot{E} , mph	53.4	53.2
Battery power at 75 mph, kW	24.4	24.3
Heat loss in battery at 75 mph 50% SOC, W	1343	1516
Average heat loss on 2 nd US06 cycle, W	1062	
Average heat loss on 3rd US06 cycle, W	1082	

Table 4.2 Comparison for calculated heat generation by Autonomie and BatPaC

It is noted that BatPaC accurately reproduced the Autonomie results for constant vehicle speed, for the standard energy demand, and for battery power at 75 mph. BatPaC slightly over estimates the heating at 50% SOC because it calculates the impedance for the parameters at that SOC. The Autonomie calculations take into account the lower values of the parameters at lower SOC and the time delay in adjusting to the ever increasing parameters with increasing SOC. The calculations in Autonomie illustrate, again, the higher rate of heat generation for high speed continuous driving than on the US06 driving cycle.

With the determination of the heat generated in the battery as a function of vehicle speed, as described above, the vehicle speed and the associated maximum temperature at the cell center are determined by the model in the following iterative procedure. The desired maximum battery temperature at steady-state is entered on line 18 and the model calculates the maximum sustained battery power, line 30, the internal battery heat generation rate, line 34, and the maximum sustained vehicle speed associated with those values, line 21. In this process, the heat generation value is transferred to the liquid cooling section beginning on line 69 or the cabin-air cooling section beginning on line 114, as determined by the type of cooling entered on line 68. In carrying out this calculation, if the temperature calculated is different from the temperature target on line 18, the vehicle speed is automatically adjusted by the program and the process is repeated until agreement is reached between the calculated temperature on line 19 and the target on line 18. The maximum temperature at the center of the cells is calculated at the end of the heat transfer calculations for the liquid-based and the cabin-air based thermal management approaches (lines 113 and 174, respectfully). As noted above, an alternative to dictating maximum cell temperature is to enter the maximum continuous vehicle speed under electric power on line 19 of the Thermal worksheet. Then the model program will override the process above and calculate the maximum cell temperature from the selected continuous vehicle speed.

The overall adequacy of the cooling is estimated on line 41 as a function of the maximum temperature of the battery at steady-state cooling as follows:

Maximum Cell Temperature	Cooling Adequacy
35 °C or less	Excellent
Over 35 °C to 40 °C	Good
Over 40 $^{\circ}$ C to 45 $^{\circ}$ C	Fair
Over 45 °C	Poor

4.2 Heating under Adiabatic Conditions

A factor to be considered in thermal management is the substantial heat capacity of the battery pack. The thermal mass of the battery evens out temperature fluctuations resulting from power bursts so that the heat dissipation system need only handle the average heat generation rate for the most extreme driving profiles the battery is likely to encounter. For large PHEV and EV batteries the heat capacity of the battery will limit the temperature rise of the centerline of the cells by distributing the heat throughout the battery. For a large EV battery with power of 120 kW and energy of 50 kWh, the temperature rise under adiabatic conditions may be only 10°C or less for a complete discharge and certainly less with a cooling system even if that system is only moderately effective. For HEV batteries, which have high power-to-energy ratios, the main effect of the heat capacity will be to smooth out temperature fluctuations.

4.3 Active Thermal Management Systems

Several heat transfer fluid types have been considered for battery pack thermal management including air from the cabin, which may be heated or cooled, ethylene-glycol/water (EG/W) solutions and dielectric liquids such as transformer fluids. Additionally, cooling plates in contact with surfaces on the cells or the modules may be cooled directly with refrigerant. Air is the least expensive, but it is less effective than the liquid coolants because of its poor conductivity, the need for large flow passages and high pumping power. Dielectric liquids are expensive, but have the advantage of being compatible with terminals and other parts at electrical potential. A 50/50 EG/W solution is inexpensive and has better properties as a coolant (thermal conductivity, heat capacity, and viscosity) than typical transformer fluids. Cooling directly with a refrigerant would eliminate the need for an intermediate heat transfer fluid and may be the most efficiency of the systems, but it is more technically complex to access, which was not attempted here. In this study we consider both thermal management approaches with heat transfer fluids of cabin air and EG/W solution. To evaluate cooling with a dielectric fluid, it would only be necessary to substitute the properties of the dielectric fluid for those of EG/W solution (lines 84-87) and to increase the cost of the thermal management system to allow for the higher cost of the heat transfer fluid. Cooling and thermal management will be used interchangeably in this report. However, we emphasize the need of the thermal management approach to allow for both cooling and heating of the battery.

For both the liquid-based and the cabin-air based thermal management approaches, we selected a general cell design that can be adapted to all of the electric-drive batteries from micro-HEVs packs to EV packs (section 2). For all types of batteries, the liquid-based design incorporates a

hermetically sealed module closure. This design can be effectively cooled by liquids and requires that only the module terminals and connectors be protected from contact with a conducting coolant, thus accommodating EG/W coolant. Aluminum heat sinks in contact with the broad surfaces of the cells transfer heat to and from the module surface (Fig. 2.4) that is in contact with the heat transfer fluid.

The enclosure does not have sufficient surface area to be cooled effectively by air. For cabin air based thermal management, heat transfers fluid flow passages are included between the cells, which add considerably to the pack volume. That design feature permits air cooling for microHEV and HEV-HP batteries. Air based approaches might also be used for some combinations of vehicle parameters and PHEV and EV battery configurations that result in low heat generation rates. However, air cooling is much less effective than liquid cooling and may not be effective in high heat generation applications (e.g. continuous all electric drive at autobahn speeds).

4.3.1 Thermal Management with Ethylene-Glycol/Water Solution

For the assumed format, the battery pack coolant is contained by the battery jacket and thus severely limits the practical pressure drop in the coolant circuit. The pack is on the suction side of the pump so the module structure assists in supporting the walls of the battery jacket. This approach is simple to calculate and appears to result in a workable design of low cost. Alternatively, cooling plates with built-in flow channels and capable of withstanding high pressure may be used with EG/W heat-transfer fluid reducing the likelihood of this electrolyte contacting cell or module terminals. Leaks in any system using an electrolyte fluid are a serious danger. For that reason alone, such fluids may have to be replaced by a dielectric fluid in future batteries.

4.3.1.1 Heat Transfer from Cell to Module Wall

As described in section 2, the cells transfer heat to the cooled walls of a hermetically sealed module with the aid of an aluminum heat conduction channel. Some of the heat is transferred through the sides of the cell to the channel and from there to the module wall. Other heat is transferred directly through the seal edge of the cell to the conduction channel flange which is in contact with the module wall. Calculation of heat transfer in this two-dimensional array through several materials is complex requiring a numerical model. The spreadsheet iterates many times in reaching a solution for the electrical and physical parameters for the battery as discussed in section 2. Each of these iterations results in a slightly different cell design. Thus, it would be impractical to imbed a numerical model directly, which may increase the total calculation time to many minutes. Instead, a software program based on the finite element method, FlexPDE 6.15 by PDE Solutions Inc., was employed to calculate heat transfer rates for 70 cell configurations. The resulting simulations were empirically correlated so that simple equations occupying a few cells in the spreadsheet could rapidly calculate the heat transfer rate with only a small error.

An important requirement for calculating heat transfer rates within the cell is to estimate the composite conductivities of the cell layers both parallel to the layers and across the layers. The resulting conductivities vary considerably with the relative thicknesses of the layers as shown in

Table 4.3, for which the results are consistent with the literature.³⁹⁻⁴³ These values for conductivities and a range of cell dimensional parameters (Table 4.4) were employed in selected arrangements for calculating heat transfer rates with the FlexPDE model for 70 representative cells that covered a broader range of variables than is needed for practical cells.

	Cell 1	Cell 2	Cell 3	Cell 4
Layer Thicknesses, microns				
Positive foil	20	20	20	20
Negative foil	12	12	12	12
Positive coating	30	75	150	220
Negative coating	40	100	200	300
Separator	20	20	20	20
Total bicell structure	212	422	772	1112
Thermal Conductivities, W/cm-K				
Aluminum	2.0	0. 2.0	0. 2.0	0. 2.0
Copper	3.8	3.8	3.8	3.8
Positive coating	0.013	0.13	0.13	0.13
Negative coating	0.013	0.13	0.13	0.13
Separator	0.0020	0.0020	0.0020	0.0020
Across layers, k_x	0.00689	0.00689	0.01045	0.0.01112
Parallel to layers, k_y	0.4127	0.4127	0.1228	0.0892

Table 4.3 Sample calculations of composite thermal conductivities of cell structures across layer and parallel to layers

Table 4.4 Range of parameter values for calculating heat transfer rates in FlexPDE model

	Parameter Levels Evaluated			
	1	2	3	4
Conductivities, W/cm-K				
Across layers, $k_x $ (a)	0.00689	0.00689	0.01045	0.0.01112
Parallel to layers, $k_y \int$	0.4127	0.4127	0.1228	0.0892
Cell edge, k_e	0.10			
Cell Dimensions, cm				
Cell thickness, <i>L</i> _{cell}	0.6	1.0	1.4	
Cell width, W	8	12	18	
Cell edge thickness, L_e	0.1			
Aluminum conductor thickness, $L_{Al}^{(b)}$	0.03	0.06	0.10	

^(a)The k_x and k_y values are calculated as in Table 4.3 and were, thus, paired together in the Flex PDE model calculations.

^(b)The total conductor thickness consists of the conductor thickness itself plus twice the thickness of the aluminum layer within the pouch material.

For each of these cells, the FlexPDE model calculated the temperature difference between the cell center and the module housing per unit of heat generation, $\Delta T/q$ (°C/W), and the fraction of

the total heat that was transferred through the edge of the cell, q_e/q . The balance was transferred through the side of the cell to the aluminum conductor, q_s . The division of the heat transfer into two routes is represented by the equation:

$$q/\Delta T = q_e/\Delta T + q_s/\Delta T \tag{4.10}$$

Estimated values for $q_e/\Delta T$ and $q_s/\Delta T$ were determined by empirical correlation of the results obtained for the calculation of the 70 cells by the FlexPDE model with the result shown in Fig. 4.1. Empirical values of these estimated values resulted in the equations:

$$q_e \Delta T = 3.917 k_y^{0.58} k_x^{-0.19} L_{cell}^{1.2} W^{0.75}$$
(4.11)

$$q_{s} \Delta T = 1628 k_{y}^{0.55} k_{x}^{0.58} L_{cell}^{-0.21} W^{0.7} L_{Al}^{0.72}$$
(4.12)



Figure 4.1 Plot comparing the estimated resistance to heat transfer from the cell center to the cooled surface of the module to that calculated by the FlexPDE model.

The average error in the estimated $\Delta T/q$ compared to the values calculated by the FlexPDE model for the 70 cases studied was 6.0% and the maximum error was 13.0%. This accuracy was deemed to be satisfactory in that for all practical battery designs, the error in estimating the difference in temperature between the cell center and the module housing, ΔT , will be only a fraction of a Celsius degree.

4.3.1.2 Heat Transfer from Module Wall to Flowing Coolant

The model directly calculates the temperature drop between the module wall and the heat transfer fluid for a set pressure drop, fluid (coolant) temperature rise, and fluid physical properties. A 50/50 ethylene glycol, deionized water (EG/W) mixture was selected based on the low cost and contemporary use in coolant systems. The default pressure drop was taken to be 10 millibar, but may be changed by the user if desired. The gap in which fluid flows is sized to maintain the target pressure drop without going below a minimum gap height of 3 mm. A coolant temperature rise of 1 °C was selected to establish a mass flow rate, but also may be changed by the user.

Calculation of the heat transfer coefficient allowed for determination of the temperature difference between the module and average coolant temperature. A schematic of the flow passageway and change in temperature profile with distance is shown below in Fig. 4.2. The outer wall of the flow passage is assumed to be perfectly insulated. The inner wall (module casing) is assumed to have a constant heat flux perpendicular to the wall. Laminar flow was assumed to simplify the calculation of the velocity profile (parabolic).



Figure 4.2 Heat transfer from the module wall to the laminar flow heat transfer fluid. The temperature profile of the fluid is shown at different lengths down the flow path.

Frequent use of dimensionless numbers was necessary to adequately correlate the numerical results into a generally useable form. We define the Reynolds, Prandlt, Graetzl, and Nusselt numbers here for completeness⁴⁴. The Reynolds number, Re, is the ratio of inertial to viscous forces. The Reynolds numbers were always less than 1000 confirming laminar flow. The Prandlt number, Pr, is the ratio of the momentum diffusivity to thermal diffusivity. The Prandlt number for the EG/W mixture is approximately 38. The Graetz number, Gz, is directly proportional to the product of the Reynolds and the Prandlt numbers. Moreover, the Gz value is inversely proportional to the distance down the fluid flow path, *l*, resulting in higher values near the start of the flow path. Finally, the Nusselt number, Nu, is the ratio of the convective to conductive heat transfer. Here u_{ave} is the average fluid velocity, d_H is the hydraulic radius (twice the flow gap), and μ is the viscosity. The heat capacity, c_p , thermal conductivity, *k*, and heat transfer coefficient, *h*, are the critical heat transfer values. The mass flow rate, *G*, and the width of the channel, *W*, are the remaining parameters.

$$\operatorname{Re} = \frac{\rho u_{ave} d_H}{\mu} \tag{4.13}$$

$$\Pr = \frac{c_p \mu}{k} \tag{4.14}$$

$$Gz = 2\frac{Gc_p d_H}{kWl} = 2\frac{d_H}{l} \operatorname{Re} \operatorname{Pr}$$
(4.15)

$$Nu = \frac{hd_H}{k}$$
(4.16)

Coupled momentum and heat transfer has been solved previously by determining a number of the eigenvalues of an analytical series solution for a vast number of various geometrical configurations related to pipe, duct, and parallel plate flow^{45,46}. We have chosen to reach the solution numerically and then fit a correlation between the Graetz number and the mean Nusselt number. The empirical form provided by Nickolay and Martin provides an accurate means of correlating the results over many orders of magnitude⁴⁷. The correlation, shown in Eq. (4.17), relates the Graetz number and the limiting solution, Nu_∞ = 5.385, to the mean Nusselt number. Then, the mean heat transfer coefficient may be directly calculated from Nu. Here *n* and *C*₁ are fitting parameters.

$$Nu = \left[\left(Nu_{\infty} \right)^{n} + \left(C_{I}Gz^{1/3} \right)^{n} \right]^{\frac{1}{n}}$$
(4.17)

The numerical model was solved with the finite element method using FlexPDE software. We note that the bulk or "cup mixing" fluid temperature in Eq. (4.18), the average temperature of the fluid normalized by the fluid velocity profile, was necessary to reach the proper values.

$$T_{avg} = \frac{1}{2u_{avg}} \int T(y)u(y)\partial y$$
(4.18)

The following important assumptions were used to reach a solution.

- 1. Flow of incompressible heat transfer fluid is laminar
- 2. Thermal diffusion is allowed up and down stream of the heat transfer (for convergence)
- 3. Boundary conditions: dT/dy = 0 at insulation; q = constant at module casing
- 4. Negligible radiative energy transfer
- 5. Steady state conditions reached

Figure 4.3 displays the temperature profile between the module casing and the insulated wall for various distances along the flow channel. The average temperature of the fluid has risen 1 °C at the end of the flow path even though the maximum and minimum temperature is separated by nearly 5 °C. The simulated change in average temperature down the length of the flow channel allows the calculation of the average heat transfer coefficient and thus Nusselt number. The correlation, Eq. (4.17), determined from various simulations conditions is shown in Fig. 4.4. An excellent fit is obtained allowing for implementation of the correlation into the design and cost model. This correlation now enables efficient and accurate calculations of the heat transfer coefficient to be made in the spreadsheet informing the user of the effectiveness of the thermal management in the design.



Figure 4.3 Temperature profile in the heat transfer fluid for various fractions of the dimensionless path length.



Figure 4.4 Correlation of model simulation results relating the Graetz number and mean Nusselt number for laminar flow between an insulated surface and the module casing.

In general, the heat transfer from the module is improved by increasing the contact area and increasing the fluid flow rate. The contact area may be increased by using cells with a higher aspect ratio. This also results in a smaller temperature gradient within the cell as discussed previously in section 4.3.1.1. Increasing the fluid flow rate is accomplished by using a lower temperature rise and/or a larger target pressure drop. The gap height may prevent a change in a single parameter from having a significant effect on the temperature drop. Physical limitations of implementing a cooling system should be considered when moving to higher flow rates and pressure drops. The user should note that raising both of these parameters will increase the cost of the battery design in ways that the model does not consider (e.g. more expensive pump, increasing structural integrity, etc).

For high speed driving or very aggressive driving cycles, the temperature difference between the surface of the cooled module surface and the bulk of the coolant may become fairly large (>10°C). This coupled with the temperature rise within the cells could result in too high cell centerline temperatures. This result can be avoided by controlling the inlet coolant temperature as a variable that is adjusted in a classic cascade automatic control system to control the module wall temperature at the desired value. Thus, the temperature rise at the center of the cells will be essentially held to that resulting from conduction within the cell and will not be greatly influenced by the temperature rise in the coolant. The use of the vehicle air-conditioning system to cool the liquid coolant allows a standard coolant temperature of 15° C, or even lower if necessary.

4.3.2 Thermal Management with Cabin Air

Air-based thermal management requires a much larger heat transfer surface area than that of liquid thermal management approach discussed in section 4.3.1. In this study the extra surface area was gained by flowing air past the faces of the individual cells as shown in Fig. 2.5. This approach greatly simplifies the calculation of heat transfer from the interior of the cells, which can be solved as a one-dimensional calculation as contrasted with the more complex two-dimensional calculation done for liquid thermal management of the outside of the module (section 4.3.1.1). The air flows in at the bottom of the long side of the battery pack, splits into many flow steams as it flows up the passages between the cells and then recombines and flows out the top of the battery pack.

4.3.2.1 Temperature Drop from Cell Center to Cell Wall at Steady State

The composite conductivity across the cell layers was calculated by dividing the total cell thickness by the sum for all layers of the conductivity of each layer divided by its thickness as illustrated in Tables 4.1 and 4.2.

The temperature drop from the center of the cell to the cell wall, ΔT_{cell} , was calculated as follows:

$$\Delta T_{cell} = \frac{\frac{q_{cell}}{2} * \frac{L_{cell}}{2}}{k_x * H_{Al} * W}$$

$$(4.19)$$
4.3.2.2 Heat Transfer from Aluminum Cell Sleeve to Flowing Air

The length of the cooled section of the cell is taken to be the same as the length of the positive electrode. The number of air flow channels is essentially calculated by dividing this length by the target width of an air flow channel plus that of one web and rounding down to an integral number. The target width was taken to be 1.0 cm, but any width between 1.0 and 2.0 cm will provide similar results for cooling. The exact width of the channels is calculated from the electrode length and the number of channels. The thickness of the channel is calculated to meet the pressure drop for the required flow rate. The required flow is that which will carry off the heat generated by the cell divided by the number of cooling channels per cell.

Because of the difficulty in moving sufficient air through the pack to carry out the heat generated in the cells, the air must be allowed to rise in temperature by about 5°C, whereas a 1°C temperature rise is sufficient for liquid-cooled packs. We also used a higher pressure drop to force the air through the pack, 0.02 bar versus 0.01 bar for liquid-cooled packs. The pressure drop for air cooling is divided between that through the channels between the cells (90%) and that in the large passages across the bottom of the pack where the air enters and across the top of the pack where the air leaves (10%). This distribution of the available pressure drop, which is achieved by sizing the thickness of the channels between the cells and those at the top and bottom of the pack, provides an even supply of air to all channels between the cells. All of the parameters discussed for managing the air flow are adjustable by the model user, but large deviations from the suggested values may adversely affect the feasibility of the design.

The heat transfer flux to establish the temperature drop from the cell sleeve wall to the bulk air stream is calculated in much the same way as described for the liquid-cooled battery packs (section 4.3.1.2). However, for the air-cooled packs, heat is transferred from both sides of the channel rather than from only one side as for the liquid-cooled modules, which have insulated jacket walls on the opposite side of the cooling channel. For transferring heat from both sides, the value of Nu_{∞} is 8.235 rather than 5.385 for use in Eq. (4.17).

4.4 Cooling and Heating Required to Maintain Pack Temperature

When parked in the sun for several hours, the internal vehicle temperature and, thus, that of the battery may become so hot that the life of the battery is reduced. To avoid this with liquid-cooled systems, the vehicle air conditioning system may be actuated intermittently to cool the battery. By allowing the temperature of the battery to fluctuate by several degrees, it is only necessary to actuate the cooling system about once per hour for a few minutes. For a set of target temperatures with a difference of 25°C and with the default insulation thickness (10 mm) and default thermal conductivity (0.00027 W/cm-K), BatPaC calculates the average cooling requirement to be about 60 W for PHEV-40 batteries. The performance coefficient of the vehicle air-conditioning system might reduce the actual energy draw to less than half that, but heating of the system outside of the battery during the hour-long downtime periods would be a counter-acting factor. The BatPaC calculates the energy required for cooling of all types of electric-drive vehicle batteries. However, most HEVs, even if cooled by liquids, may not have electrically driven air-conditioning units and some other method might be needed to avoid very high battery temperatures during parking such as thicker insulation and fan cooling during parking. Batteries

for any type of electric vehicle that are cooled by air would require similar measures or some method of connecting the air cooling to an electrically-driven air-conditioning system or dedicated refrigeration system. The additional cost for any special system to deal with high ambient temperature that is not a part of the system to remove heat generated within the battery has not been provided in the BatPaC model.

If the battery is to deliver full power at startup, it must be at a temperature of at least 5°C. This minimum temperature can be maintained by heaters and circulation of the coolant, whether air or liquid. The BatPaC calculates the amount of power required to maintain the battery temperature for any set of battery and ambient temperatures. PHEV-40 batteries would require about 50 W of heat to maintain the battery temperature at 20°C above that of the ambient under steady-state conditions. During recharging this should be easily done for 20 hours at a cost of \$0.10 with an energy cost of \$0.10 per kWh. If the vehicle is not at a source of power for recharging, limited energy (say 1-2 kWh) can be drawn from the battery (if not blocked by a switch actuated by the driver) and then automatically shut off after maintaining the battery temperature for one to three days, depending on the ambient temperature.

4.5 Heat-up from Cold Ambient Conditions

All of the batteries for the various types of electric-drive vehicles will occasionally be exposed to very cold temperatures, which will require special heat-up procedures. All but the EV batteries can be heated with the aid of the engine. This can be done with electric heaters operated from power taken from the generator or from glycol solution from the engine cooling system. If the latter, it might be prudent to isolate the engine coolant from the battery coolant by means of a plastic heat exchanger.

Another method of heating the battery is by means of the electric heaters that should be available for maintaining the battery temperature (section 4.4). BatPaC calculates the amount of heat needed and the time required with suitable heater power. For PHEV-40 batteries, about 15 minutes is required with 2-kW heaters. This method of heating will be slower than with the engine coolant and even the latter would result in some delay before the battery is capable of full power.

To avoid delay in starting vehicles from a cold startup, the driver could initiate heating by means of a remote device, even a telephone. By this means, heating could be initiated either from heat drawn from the engine or electric heaters. Remote initiation of heat-up would be especially important for an EV away from a charging station in that no engine is available to assist heating and the large size of the battery would result in a long heating period with electric heaters. The BatPaC estimates the time under these conditions.

5. Modeling of Battery Pack Manufacturing Cost

5.1 Approach

The manufactured cost of a battery pack is calculated with input from the design information generated in modeling the cell and battery pack performance. The design modeling determines the annual materials and purchased items requirements. The manufacturing cost is then added to these materials costs, along with a warranty cost, to reach the unit cost of a single battery pack. The manufacturing costs for the designed battery are scaled from a baseline plant. The baseline plant was designed for a battery of intermediate size and production scale so as to establish a center-point for other designs. The baseline plant accounts for the size, speed, number of units, direct labor, and depreciation of the capital cost for each processing step. These costs are adjusted to meet the requirements for a plant producing the battery under study. The process expenses are summed with the additional costs of operating the manufacturing facility. These costs include launch costs, working capital, variable overhead, general, sales, administration (GSA), research and development, depreciation, and profit. Additionally, the costs for the thermal management, battery management system, and disconnects have been estimated to provide the total cost to the OEM for the integrated battery pack.

In this analysis, all costs are evaluated for 2020 when large battery manufacturing plants are built. All dollar values are brought back to 2010 with allowance for inflation. In other words, all costs and prices are in 2010 dollars. Some materials and battery manufacturing costs are lower than recent values, where we judged that processing improvements for high volume production of materials would reduce costs.

The baseline manufacturing plant was calculated for an annual production rate of 100,000 batteries. The cost model accounts for different scales of manufacture by recalculating the costs of each individual step in the manufacturing process. The changes in capital and operating costs will change the calculated unit cost of the battery pack. The parameters were determined to provide reasonable estimates for manufacturing rates of 20-500 % of the baseline rate. Thus, for a plant that is far different in size from the baseline plant, for instance a pilot plant having an annual production of only 5,000 battery packs per year, the estimate from this study would be expected to be less accurate than if determined in a study dedicated to that purpose.

To simplify the cost calculations, it was assumed that all hardware items for the cells, modules and battery will be purchased from a vendor specializing in similar products. The costs for these items were estimated to be a fixed value plus an additional value proportional to the weight of the item, which is calculated during the battery design. In mature manufacturing plants in 2020, toward which this study is directed, some items which are assumed to be purchased in this study might actually be internally manufactured from raw materials. This would increase the number of processing steps needed in our manufacturing simulation and thus complicate the cost calculations. Assuming that some parts would be purchased if they would actually be produced from raw materials would tend to underestimate capital and labor costs and overestimate purchased items expenses. However, the net effect would be a very small change to the overall unit cost of the battery pack.

5.2 Materials Costs, Purchased Items, and Pack Integration

The end battery pack cost depends significantly on the cost of both the active and inactive materials that compose the design. In this subsection, the assumed material costs and the rationale behind them are presented. We provide for means to scale the materials cost with production volume using the same method used for processing rates as discussed in section 5.4. In general, the materials costs will be largely insensitive to production volume since we have assumed a high volume market already exists. Only the negative and positive electrode active materials are assumed to have a minor benefit for larger scales of production. While we state suggested materials costs and sensitivity to production scale, the users of the cost model may enter any value that they desire.

5.2.1 Battery Specific Materials Cost

The largest contributions to the materials cost of the battery are from the following components: positive and negative electrode active material, separator, electrolyte, and current collector foils. The choice of the materials often defines the size and performance of the battery as well as the cost. Many different variations of materials are possible in the Li-ion family of chemistries. However, we have chosen to focus on the different available positive electrode materials with less attention on the negative electrode. This reflects the current research and manufacturing activities. The separator and the electrolyte are also both active areas of development. However, the following battery designs are based on a single electrolyte and separator combination. Including the cost and effect of additives and enhanced separators is beyond the scope of this work. The user is always able to modify the dimensions, cost, and ASI that may be required to account for changes in these materials.

The price of specific battery materials is of some debate. The values presented in Table 5.1 compare our suggested costs to those reported recently in the open literature. Our values, as well as the others in the table, are derived from conversations with material, cell, and original equipment manufacturers. The sources are commonly anonymous and the accuracy of the values is generally unknown. We present the comparison of published values so that the user of the cost model may appreciate the accepted range of values for commonly used materials.

5.2.1.2 Positive Electrode Active Materials

The cost of positive electrode materials is driven to a large extent by the cost of the raw materials from which it is made. The archetype Li-ion positive electrode material, lithium cobalt oxide (LCO), was the original material commercialized in Li-ion batteries for consumer electronics. LCO has many excellent characteristics but is not considered a viable choice for use in Li-ion batteries for automotive applications. One of the largest drawbacks of LCO, other than safety concerns, is the high and volatile cost of the cobalt. While tolerable in the consumer electronics market, the cost is too high for use in an automobile battery. Many other materials are in a commercially viable state of development and are currently utilized in Li-ion batteries produced today (Table 5.1) such as lithium manganese spinel oxide (LMO) and lithium nickel manganese cobalt oxide (NMC).^{3,6} The relative advantages and disadvantages of each material will not be discussed here.

3,6
and separator.
electrolyte,
anodes,
for cathodes,
costs
of stated
Details c
Table 5.1

				ANL	THAX**	CARB
Material	Chemistry	Abbreviation	unit	2010	2010	2007
Manganese spinel cathode	$Li_{1.06}Mn_{1.94-x}M_xO_4$	LMO	\$/kg	10	12 - 16 - 20	8 - 10
5V spinel cathode*	$LiNi_{0.5}Mn_{1.5}O_4$	LNMO	\$/kg	21	I	I
Phospholivine cathode	$LiFePO_4$	LFP	\$/kg	20	15 - 20 - 25	16 - 20
Layered oxide cathode*	$LiNi_{0.80}Co_{0.15}Al_{0.05}O_2$	NCA	\$/kg	33 // 37	34 - 40 - 54	28 - 30
Layered oxide cathode*	$Li_{1.05}(Ni_{1/3}Mn_{1/3}Co_{1/3})_{0.95}O_2$	NMC-333	\$/kg	31 // 38	40 - 45 -53	22 - 25
Layered oxide cathode*	$Li_{1.05}(Ni_{4/9}Mn_{4/9}Co_{1/9})_{0.95}O_2$	NMC-441	\$/kg	26 // 29	I	I
Li & Mn rich layered cathode*	$xLi_2MnO_3{\boldsymbol{\cdot}}(1{\boldsymbol{\cdot}} x)LiNi_yMn_zCo_{1{\boldsymbol{\cdot}} y{\boldsymbol{\cdot}} z}O_2$	LMR-NMC	\$/kg	22-25 // 22-29	24 - 31 - 39	I
Layered oxide cathode*	LiCoO ₂	LCO	\$/kg	36 // 59	I	30 - 40
Synthetic graphite anode	C_6	Gr	\$/kg	19	17 - 20 - 23	I
Natural graphite anode	C ₆	Gr-Nat	\$/kg	10	I	I
Titanate spinel anode	Li ₄ Ti ₅ O ₁₂	LTO	\$/kg	12	9 - 10 - 12	I
Electrolyte	1.2 M LiPF_6 in EC:EMC		\$/kg	18	18.5 - 21.5 - 24.5	I
Separator	PP/PE/PP		m^2	2	1 - 2.5 - 2.9	I
Current collector foil	Copper		\$/m ²	1.80	I	I
Current collector foil	Aluminum		\$/m ²	0.80	I	I
* The cost of cathode materials usin	g co-precipitation of Ni, Mn, and/or Co is b	ased off of a correl	ation. Co	balt \$44/kg vs. \$81/k	g separated by "//"	
** Cost represent range of values po	ossible					

The amount of cobalt and nickel, as well as ease of manufacture, controls the end price for a positive electrode material. For example, the NMC-441 is less expensive than the NMC-333 as the cobalt quantity is significantly reduced. The market prices for cobalt and nickel metals vary dramatically from year to year. Reducing the quantities of these materials in the positive electrode will reduce the total price and price volatility. Researchers at TIAX LLC have treated this variation and shown the significant effect on end battery cost.¹⁰ The average traded metal prices and the 95% confidence intervals for the last 25 years is 44.4 ± 18.3 \$/kg and 14.9 ± 7.6 \$/kg for cobalt and nickel respectively. These numbers are based on historical prices for the metals as collected by the United States Geological Survey (USGS).⁴⁸ The metal prices are indicators for how the intercalation material cost will relate when compared to one another. The fact these materials are not earth abundant means they will not benefit as much as other materials from increased scales of production.

We employ the relationship in Equation 5.1 to systematically calculate the cost of the transition metal based spinel and layered compounds. The final cost, C, of the lithiated oxide depends on the baseline cost, C_0 , and the contributions of the lithium and transition metal raw materials, C_i . The molar stoichiometry, x_i , is transformed to a mass basis with the molecular weight of the raw material, MW_i, and the final product, MW. The baseline cost is the sum of the cost for processing, additional raw materials, and profit margin associated with the manufacture of the materials. We assume a baseline cost of \$7/kg for single metal containing oxides (LMO and LCO) and \$16/kg for the co-precipitated metal oxides such as NMC-333 and NMC-441. NCA is known to have a slightly lower yield and requires additional raw materials resulting in an assumed $C_0 =$ \$20/kg. The costs for Li, Ni, Mn, and Co are taken to be 0.22, 0.87, 0.15, and 2.6 \$/mol respectively. The historical average metal prices for Ni and Co are used recognizing that these values will fluctuate over time. Aluminum is assumed to be similar in cost to manganese for these calculations. One may directly translate these numbers to raw materials costs resulting in \$6/kg for Li₂CO₃, \$5.6/kg for NiSO₄, \$16.9/kg for CoSO₄, and \$1/kg for MnSO₄. Calculations are also shown in Table 5.1 using \$4.8/mol (\$81/kg) for cobalt as a demonstration of the upper 95% confidence interval cobalt price on the end material cost.

$$C\left(\frac{\$}{kg}\right) = C_0 + \frac{1}{MW} \sum_i x_i C_i MW_i$$
(5.1)

In general, earth abundant elements should be the dominate transition metals used if a low cost positive electrode is desired. Both iron and manganese are abundant and inexpensive transition metals for intercalation materials. Comparison of the iron phosphate, LFP, to manganese spinel, LMO, reveals how processing costs contribute to the end price of a material. LMO is relatively easy to manufacture. In contrast, LFP requires a reducing atmosphere and a carbon coating step to reach the end product. The increased complexity in the manufacturing process is realized in the price. However, one could argue that the manufacturing cost will decrease with increased knowledge from larger scales of production.

5.2.1.2 Negative Electrode Active Materials

While several negative electrode materials exist for Li-ion batteries, carbon materials in the form of graphite and/or hard carbon are still used in the vast majority of commercial cells. Graphite

offers the greatest energy density while hard carbon is said to enable high rate capability with decreased risk of lithium plating (an undesired side reaction) during high charge rates. We have chosen synthetic graphite as a generic carbon electrode in our model. Significant differences in cost and performance will exist between synthetic, natural, and coated-natural graphite. The method of production and necessary heat-treatment will control the end cost. Many manufacturers use a proprietary blend of natural and synthetic graphite and/or hard carbon in the negative electrode of their cells. The user of the model may feel free to vary the price depending on the application of interest.

The lithium titanate electrode, LTO, offers an interesting option compared to graphite. Unlike graphite, LTO operates within the stability window of the electrolyte. The higher electrode potential, 1.5 V vs Li, dramatically reduces or eliminates the formation of the solid electrolyte interphase (SEI). As a result, nanoparticle-based LTO may be implemented without concerns of increased side reactions with the electrolyte. The reduced nanoparticle dimensions increase the available surface area for reaction while simultaneously shortening the diffusion length. Both of these factors combined with the lack of SEI dramatically reduce the impedance of the electrode.

5.2.1.3 Electrolyte and Separator

The electrolyte used in this model is based on a lithium hexafluorophosphate salt, LiPF₆, dissolved in a carbonate based solvent system. The carbonate solvent system is a blend of ethylene carbonate, EC, and a linear carbonate such as ethyl methyl carbonate, EMC, or dimethyl carbonate, DMC. Other chemical additives may be used to limit the capacity and power fade of the battery over time. Polymers may be added to the electrolyte as either a minor or major component. This is not discussed in any further detail in this work. The price of 18 k, about 22 /L, is only for the base electrolyte (i.e. no additional additives).

The separator is typically a porous membrane based on polypropylene (PP) and sometimes includes a polyethylene (PE) middle layer. PP and PE are very inexpensive raw materials and thus the suggested cost of $2/m^2$ is in large part due to the manufacturing process required to form the porous network in the membrane. As competition and scale of manufacture increase, the prices of the separator may fall closer to $1/m^2$. However, the cost of improved technology may offset some of this cost reduction, so we have retained our cost estimate of $2/m^2$. As safety is a major concern for Li-ion batteries, the separator plays a key role in isolating the oxidant from the fuel. If the two charged electrodes contact each other (short), then a run-away reaction is possible. Separators have been designed to "shut-down" or melt at key temperatures. The middle PE layer is the shut-down feature in our proposed separator. Ceramic coatings have also been used to ensure structural integrity. Many other approaches are being developed to increase the safety of Li-ion batteries. The user of the cost model should account for the specific separator technology in the price and dimensions (thickness and porosity) of the separator as needed.

5.2.1.3 Current Collector Foils

The current collector foils are based on copper metal for the negative electrode and aluminum for the positive electrode. However, the LTO anode material, because of its high voltage relative to lithium, enables the use of aluminum as the negative electrode current collector. The price of these foils is based on raw materials and manufacturing costs. The aluminum foil is produced by rolling of thicker stock foils into thinner and thinner sheets. On the other hand, copper foil is more likely to be produced through an electrodeposition process. The foils are 12 microns and 20 microns thick for the copper and aluminum current collectors respectively. The foils used in batteries have additional requirements beyond the cheapest product available. Surface treatments are often necessary to promote adhesion of the composite electrode to the foil surface. In addition, alloying of the foil may be necessary to achieve the required material properties for long life.

The raw material contributions to the foil price will vary with the volatility of the market price for the metals. Figure 5.1 displays the metal ingot price contribution on a $/m^2$ basis. These numbers are based on historical prices for the metals as collected by the USGS.⁴⁸ The values for both aluminum and copper tend to vary significantly over the time period examined. The price for copper is more volatile and always more expensive than aluminum. Analysis of Figure 5.1 reminds the user of the cost model that cost quotes are only valid for a short period. As the market price for raw materials changes, so will the price for the finished product.

Conversations with manufacturers and suppliers lead us to take a price of 1.80 and $0.80 \text{ }^2/\text{m}^2$ for battery grade copper and aluminum foil respectively. We point out that the current metal ingot price is only a small contribution to the end foil price being about 16 % of the aluminum foil price and 23 % of the copper foil price. Thus, a doubling of the ingot prices would only moderately increase the foil prices.



Figure 5.1 Metal ingot cost contribution to the current collector foils over a 20 year period. The average for that period is also shown. All costs are in 2010 US\$.

5.2.1.4 Additional Electrode Components

The binder and conductive additive in the positive and negative electrodes add a small but real cost to the battery. The conductive additive, more common for the positive electrode, was priced at 6.80 \$/kg for a high purity and moderate surface area carbon black material. The binder, perhaps PVDF or CMC based, is assumed to be 10 \$/kg. The *N*-Methyl-2-pyrrolidone (NMP) solvent for the PVDF binder is estimated to be 3.20 \$/kg. Most of the NMP is recovered after evaporation and recycled as discussed in section 5.3.3. Only the small amount lost in processing need be replaced. No cost is assumed for water used in the electrode slurry processing.

5.2.2 Purchased Items Cost

Table 5.2 lists the purchased items for the cell module and battery jacket. The cost of a SOC controller for each cell, or group of parallel cells, is \$2.50 plus a small factor for the cell capacity (Ah), which allows for higher cell balancing currents for larger cells. The other component costs include a fixed amount plus an additional factor, which is proportional to their mass, m_i . The cell negative terminal and parallel cell group connection are both made from nickel plated copper sheet and thus have the same cost equation. The costs shown for the terminals include an allotment for isolation tape that is necessary to protect the electrical connection. The bus bar is a fixed cost and is only charged if a single row of modules is used. A single row of modules requires a bus bar in order to locate the positive and negative terminals at the same end of the battery.

Component, <i>i</i>	Cost Equation, \$/unit	Cost per unit
SOC controller	2.50 + 0.01C	cell or parallel cell group
Cell positive terminal	$0.25 + 4m_i$	cell
Cell negative terminal	$0.25 + 6m_i$	cell
Cell container	$0.20 + 3m_i$	cell
Aluminum heat conductor	$0.10 + 4m_i$	cell
Parallel cell group connection	$0.25 + 6m_i$	parallel cell group
Module terminals	$0.75 + 5m_i$	module
Balance of module (casing)	$1.00 + 3m_i$	module
Module interconnect	$1.00 + 5m_i$	module
Battery terminals	$15.00 + 0.02I_{total}$	battery pack
Bus bar for one module row	20.00	battery pack
Battery jacket	$30.00 + 7m_i$	battery pack

Table 5.2 Cost equations for purchased items

5.2.3 Pack Integration Cost

Various additional components and thus cost are necessary to integrate the battery into the electric drive system, which adds cost. While it is not clear what should and should not constitute the cost of the "battery pack," we present these additional items in Table 5.3 in an attempt to be complete. After all, the cost of the entire system is of interest to the final consumer of the product. The calculated total battery cost to the OEM may be directly compared to battery pack level goals as set by the U.S. Advanced Battery Consortium. This category in BatPaC includes

the battery pack as well as the battery management system and disconnects. However, thermal management additions outside of the pack are only included in the total system cost to the OEM. The general conclusion is that the pack integration costs have the largest consequence for the smallest batteries. The worst case is perhaps that of a small PHEV10 battery. The integration costs of a PHEV10 battery carry the burden of charging from the grid, but provide only a modest electric drive benefit. The fixed cost of pack integration may amount to 25 % of the battery pack total even without considering the costs of additional powertrain components. Clearly, understanding the entire cost of the electric drive system is of importance to evaluating the true value of the electrified vehicle to a consumer.

Battery Pack Integration	System		PHEV
	MicroHEV	HEV-HP	and EV
Current and voltage sensing, \$	40	70	100
Module controls, \$/module	10	10	20
Auto. battery disconnect, \$	50	70	200
Manual disconnect per pack, \$	15	15	15
Additional for parallel modules and packs, \$		per string	100
Thermal Controls			PHEV
	MicroHEV	HEV-HP	and EV
Baseline thermal system*, \$	30	80	120
Additions to AC system**, \$/kW	40	40	40
Heating system**, \$/kW/pack	20	20	20
Additional for multiple packs, \$/a	dditional pac	:k	100
*\$60 additional for each added pa	ack		
**No charge for cabin air cooling			

Table 5.3 Costs to integrate battery pack into vehicle drivetrain. \$/kW numbers reflect maximum kW of cooling or heating required.

5.2.3.1 Battery Management System

The battery management system (BMS), in our assumed battery design, integrates the modules and battery into the overall electric drive system. The BMS includes measurement and control features such as the following:

- Measurement of battery pack current and voltage
- Balancing of the module voltages (cell balancing done within module)
- Estimation of battery pack state-of-charge (SOC) and state-of-health (SOH)
- Estimation of module SOC and SOH
- Monitoring and signaling of battery thermal management

The cost of the BMS will scale with magnitude of battery current and with the need to charge from the electrical grid. Therefore the PHEV and EV batteries will have a higher burden from the BMS. The micro-HEV is assumed to have less complicated management and thus less cost than the HEV-HP.

5.2.3.2 Manual and Automatic Disconnects

The manual and automatic disconnects integrate a high-level of safety and electrical management into the electric drive system. The manual disconnect breaks the current flow pathway from the high-voltage terminals to the outer system allowing for the safe service of the vehicle and battery pack. This disconnect is designed to be operated when the electrical system is de-energized. The automatic disconnect is much more complex. This unit contains the connections for the highvoltage system to the rest of the vehicle's electrical system: drivetrain, grid charging (if applicable) and accessories (high and low voltage). Fuses are present as a hard-wired safety device to prevent unusually large current spikes from damaging the battery or drivetrain. Multiple contactors are used to appropriately channel electrical current depending upon normal operation or grid-charging. Engaging the contactors requires that multiple safety interlocks are established including isolation of the high voltage bus from the vehicle chassis and an inertia based sensor (crash protection). Finally, a small circuit is provided to prevent arcing of the current across the high-voltage contactor when the high-voltage circuit is closed.

The relative cost of the automatic disconnect amongst the various battery designs is driven by the pack voltage, maximum battery current, and the need for charging from the grid. The voltage of the pack has a significant effect if a 42 V micro-HEV pack is considered. For this system, electrical safety regulations allow a less complicated system to be used. Requiring higher battery currents generally increases the cost of electronics and conductors. The additional complications arising from grid-charging adds a significant additional cost to the PHEV and EV systems. It is unclear to the authors at this time what other factors may enable a lower burden of external safety controls. These additional costs in the automatic disconnect unit have the most pronounced effect on the cost of smaller batteries, as the burden amounts to a significant fraction of the total cost.

5.2.3.3 Balance of Thermal Management System

The thermal management of the battery is crucial to meeting the life and safety requirements of transportation applications. The complexity of this system must be minimized to reduce the cost and size burden on the vehicle. A single refrigerant compressor is used for both the cabin air and the battery cooling applications. Likewise, the same radiator and fan as the cabin cooling will also be used for the battery cooling refrigerant. Most OEMs appear to use an electric compressor for all full-HEVs (HEV-HP) and PHEVs/EVs that are liquid cooled. The incremental cost for an electric compressor at high volume in year 2020 will likely be \$200-300 more than the commonly used \$100 belt driven compressor. We do not include this incremental cost in our thermal management system cost; however, we state it here for completeness. The additional cost to the compressor for the battery cooling capacity is insignificant compared to burden of transitioning to the electric compressor. Experts in the field have informed us that the electric motor and high-voltage invertor are the largest contribution to the incremental cost of the electric compressor.

An expansion valve on the refrigerant line and a heat exchanger (chiller) transfers the thermal energy from the heat transfer fluid to the refrigerant loop. A 50/50 DI-water/ethylene-glycol solution is selected as the heat transfer fluid. The assumed battery design has the heat transfer

fluid pumped over the module casing to transfer heat from and to the cells. The battery may be heated by either a positive thermal coefficient (PTC) or flexible mat heater. The PTC heater would directly raise the temperature of the heat transfer fluid in a reservoir while the matt heater would be placed under the battery jacket insulation.

The PHEV and EV batteries will likely have both active and passive thermal management modes requiring some additional monitoring and an electrically actuated valve. We have assumed decreasing cooling costs for the HEV-HP and microHEV systems without explicitly dictating where the savings originate. In general, one would expect smaller batteries to have a less complicated control system, lower flow rates and possibly even direct cooling of the cells with the evaporator.

5.3 Baseline Manufacturing Plant

The model's baseline plant is designed to produce 100,000 NCA-Gr baseline battery packs per year. The baseline battery pack produced by the plant has sixty, 40-Ah capacity cells, providing a total pack power of 50 kW and total energy of 8.7 kWh. The battery will power 24 miles of vehicle travel at 70% of the pack energy and 250 Wh/mile. The schematic diagram of the plant (Fig. 5.2) is designed to illustrate the flow of materials through the plant and the relative floor areas for the processing steps rather than representing a realistic plant layout. The overall manufacturing rate of 100,000 battery packs per year is achieved by operating for three shifts at the equivalent of 300 days per year of fully effective production. There will be more than 300 days of operation, but all days will have less than 100% effectiveness. The exceptions to three-shift operation are the Receiving and Shipping sections, which are active for only two shifts per day. The cost factors for the individual manufacturing steps in the baseline plant are summarized in Table 5.4 and discussed in detail in the sections that follow. Most of the operations are carried out with normal factory atmosphere, but the cell assembly process steps are completed in a dry room atmosphere.



The areas in this diagram for each processing step are approximately proportional to the estimated plant areas in the baseline plant.

Figure 5.2 Baseline lithium-ion battery manufacturing plant schematic diagram. Manufacturing rate: 100,000 NCA-Gr battery packs per year, 50-kW pack power, 40-Ah capacity, 60 cells per battery. Operating year: 300 days with three 8-h shifts (two shifts for receiving and shipping)

5.3.1 Receiving and Shipping

These operations incorporate the moving equipment and storage facilities common to any such factory facilities. The Receiving section handles slightly less than 6,000,000 kg of materials per year and also has facilities to handle and store some of the electrode materials in a dry atmosphere. The Shipping section is required to enclose the battery packs in crates, which requires some automated equipment and more labor than is required for Receiving. Shipping also handles about 400,000 kg of scrap each year, which is broken down and prepared for shipping in the Rejected Cell and Scrap Recycle section. The estimated resources needed for the Receiving and Shipping sections are shown in the table below.

	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
Receiving	870,000 kWh/y	3 per shift	3.60 mil\$ total	900
Off-loading			0.60	
Moving			1.20	
Storage			1.80	
Shipping	870,000 kWh/y	6 per shift	5.0 mil\$ total	900

*Total cost including installation

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100,000	Battery Pa	cks per Year, 50-k	W Battery F	Power, 40-/	Ah Capacit	y, 60 Cells	per Battery		
Operati	ing year. suu	uays with three o-r	I) STILLS INOL		er day ior re	ceiving and	(Buiddius)	i	
				Direct Labo	-	Cap. Eq	uipment	Plant	Area
			Workers/	Baseline	Scale	Baseline	Scale	Baseline	Scale
	Annual B	aseline Rate (R _o)	Shift	Hours/yr	Factor, p	\$millions	Factor, p	Area, m²	Factor, p
Receiving (two-shift operation)	869,416	kWh energy	e	14,400	0.4	3.60	0.6	006	0.5
Materials preparation									
Positive electrode	1,712,524	kg active material	2	14,400	0.5	2.00	0.7	600	0.6
Negative electrode	1,060,374	kg active material	7	14,400	0.5	2.00	0.7	600	0.6
Electrode coating				ı					
Positive electrode	8,209,039	m ² cell area	4	28,800	0.5	8.00	0.8 (0.2)*	750	0.8
Negative electrode	8,209,039	m ² cell area	4	28,800	0.5	8.00	0.8 (0.2)*	750	0.8
Solvent recovery	2,309,021	kg NMP	7	14,400	0.4	3.00	0.6	225	0.6
Calendering									
Positive electrode	8,209,039	m ² cell area	7	14,400	0.5	1.00	0.7	225	0.6
Negative electrode	8,209,039	m ² cell area	-	7,200	0.5	1.00	0.7	225	0.6
Materials handling [#]	8,209,039	m ² cell area	4	28,800	0.7	1.50	0.7	006	0.6
Electrode slitting	8,209,039	m ² cell area	4	28,800	0.5	2.00	0.7	300	0.6
Vacuum drying	8,209,039	m ² cell area	7	14,400	0.5	1.60	0.7	300	0.6
Control laboratory	869,416	kWh energy	4	28,800	0.5	1.50	0.7	300	0.6
Cell Assembly in Dry Room									
Cell stacking	6,315,789	total cells	5	36,000	0.7	4.00	0.8 (0.3)**	600	0.8
Current collector welding	6,315,789	total cells	5	36,000	0.7	4.00	0.8	600	0.8
Enclosing cell in container	6,315,789	total cells	ო	21,600	0.5	3.00	0.7	600	0.6
Electrolyte filling, and cell sealing	6,315,789	total cells	5	36,000	0.5	5.00	0.7	006	0.6
Dry room control and air locks	2,000	m ² operating area [#]	7	14,400	0.4	20.00	0.6	100	0.4
Formation cycling	6,315,789	total cells	ω	57,600	0.7	30.00	0.8 (0.3)**	2,200	0.8
Final cell sealing	6,315,789	total cells	7	14,400	0.5	2.00	0.7	450	0.6
Charge retention testing	6,315,789	total cells	ო	21,600	0.4	4.75	0.7	006	0.6
Module assembly	6,000,000	finished cells	9	43,200	0.5	6.00	0.7	600	0.6
Battery pack assembly and testing	100,000	battery packs	9	43,200	0.5	6.00	0.7 (0.3)***	006	0.6
Rejected cell and scrap recycle	6,315,789	total cells	5	36,000	0.7	2.50	0.7	600	0.6
Shipping (two-shift operation)	869,416	kWh energy	6	28,800	0.5	5.00	0.7	006	0.6
Total			06	626,400		127.45		15,425	
#One-third of the space for materials	handling is w	ithin the dry room.							
*The baseline cost for capital equipm	nent for electr	ode coating, Co, is ba	sed on the ev	aporation of t	he baseline a	innual solvent	t weight (Rs _o)	of 2,309,000	kg for
the positive electrode and 1,527,000	kg for the neg	jative electrode. For t	patteries requ	iring different	solvent evap	oration rates	Rs, the cost	is multiplied b	Ž
(annual solvent rate (Rs)/baseline rat	te (Rs _o)) raise	d to the 0.2 power. Th	hus, Cost =	Co*(R/Ro)0.	8*(Rs/Rs _o)⁄0.	5			
**The baseline costs of the capital er	quipment for a	ell stacking and form	ation cycling	is for 40-Ah c	ells. To corr	ect the baseli	ine cost (C _o)	for cells of dif	ferent
capacity, the cost is multiplied by the	he capacity ra	itio, (Cap)/ 40 Ah, rais	ed to the 0.3	power. Thus	t, Cost = C _o *	(R/Ro) 0.8*(C	ap/40)^0.3.		
***The baseline cost of the capital ec	quipment for b	attery assembly is for	a battery wit	h four module	es. To correc	t the baseline	e cost to a be	ittery with a d	fferent
number of modules (Mod), the cost	is multiplied t	y the ratio of the num	ber of module	es, (Mod)/4 ra	aised to the 0	.3 power. Co	$st = C_o^*(R/R)$	o)^0.7*(Mod/4	M.3.

5.3.2 Electrode Materials Preparation and Delivery to Coating

The electrode materials, which consist of active material, carbon (if necessary), binder and binder solvent, are well mixed in small batches in portable tanks. At the design production rate in the baseline plant, each shift requires three tanks each holding about 1000 liters of positive electrode material mix and three tanks each holding about 900 liters of negative electrode material mix. The section must be capable of exceeding this design rate of production by at least 25% to catch up in case of unscheduled downtime in Materials Preparation or in some of the section immediately following that section. The tanks of prepared materials are moved to the Coating section and pressurized to push the coating paste into the coating mechanism. The estimated resources needed are the following:

Materials Prep.	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
Positive	1,710,000 kg/y	2 per shift	2.0 mil\$ total	600
Materials	active material			
Storage tanks			1.00 mil\$	
Mixing tanks			0.50	
Moving equip.			0.50	
Negative	1,210,000 kg/y	2 per shift	2.0 mil\$ total	600
Materials	active material			

*Total cost including installation

5.3.3 Electrode Coating on Current-Collector Foil

The positive and negative electrode structures are formed by coating both sides of the current collector foil. In the baseline plant, the coating lines are 1.5 meter wide continuous roll-to-roll coating processes carried out at a line speed of 10 m/min. The first set of coating and drying stations coats one side of the current collector foil, drives off the solvent in a heated oven, and turns the foil over while transferring it to a second set of stations. The second set of coating and drying stations applies and dries the remaining coating before the coated foil is wound into a large roll at the end of the line. An advanced alternative would be to run the foil directly into the calendering process. The negative and positive coating lines are very similar. However, some of the negative material is coated only on one side to provide the electrodes at the end of the cell stacks. For the baseline plant, a total of 8,170,000 m^2/y of coating (annual cell area) is required for the positive electrode (slightly more for the negative electrode), which allows for the 5% loss of cells expected to fail testing and inspection. A larger area of foil than the coated area must be fed to the coaters to allow for the part of the foil that is not coated so as to provide tabs for welding to the terminals (about 10%) and to allow for trimming losses during electrode slitting (8%). Also, about 30% excess coating capacity must be provided to allow for unscheduled downtime. Only one coating line is needed for each electrode type to meet these needs. If one coating line breaks down, the other coating line may change over temporarily to coat the other electrode material.

The oven sections of the coating line are designed to dry coatings about 100 microns thick at the coating speed of 10 m/min. A thicker coating will require longer ovens at additional capital cost which is provided in the adjustment of costs discussed in section 5.4. For the same annual area

throughput, a coating line that coats both sides with a 300-micron coating would cost \$9,500,000 rather than the \$8,000,000 cost for the 100-micron coater. The binder solvent for the positive electrode in the baseline plant is NMP, which must be recovered by condensation and recycled. About 0.5% of the binder solvent is combusted with a thermal oxidizer and must be replaced. For the negative electrode the binder is water, which need not be recovered. The estimated resources to meet these needs are the following:

Electrode Coating	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
Positive Electrode	8,170,000	4 per shift	8.0 mil\$ total	750
	m ² /y cell area			
Uncoated area			18%	
Width of coater			1.5 m	
Coating speed			10 m/min	
Number of coaters			One	
Maximum rate			$13,000,000 \text{ m}^2/\text{y}$	
Excess capacity			30%	
Negative Electrode	8,170,000	4 per shift	8.0 mil\$ total	750
	m ² /y cell area			
Solvent Recovery &	1,527,000 kg	2 per shift	3.0 mil\$ total	225
Oxidation	NMP/y			

*Total cost including installation

5.3.4 Calendering

The materials leaving the coating lines may be stored on large rolls (see next section). However, typically the materials leaving the coaters would go directly to the calendering process in which the coatings are compressed by rolling to meet the specified void volume fraction, which will later be filled with electrolyte. The calendering equipment must match the output of the coating equipment producing 8,170,000 m²/y of cell area with a maximum rate of 13,000,000 m² of foil per year to meet contingencies as in coating. We estimate three workers are necessary to collectively operate the two pieces of equipment. The estimated resources to meet these needs are the following:

Calendering	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
Positive Electrode	8,170,000	2 per shift	1.0 mil\$ total	225
	m ² /y cell area	-		
Negative Electrode	8,170,000	1 per shift	1.0 mil\$ total	225
-	m ² /y cell area			

*Total cost including installation

5.3.5 Inter-Process Materials Handling

For all processes (Fig. 5.2), work in progress must be transported and occasionally stored to permit nearly-continuous operation of the equipment. Storage areas must be provided both inside and outside of the dry room. Raw materials must also be moved to the processing sites, which for those in the dry room means through a separate air lock for materials transfer. One-third of the

total space for Inter-Process Materials Handling is within the dry-room for the baseline plant and also for all other plants. The estimated resources to meet these needs are the following:

Materials Handling	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
	8,170,000	4 per shift	1.5 mil\$ total	900
	III / y cell alea			

*Total cost including installation

5.3.6 Electrode Slitting

The coated electrode foils are slit into strips between the coated sections and then into individual electrodes as shown in Fig. 2.3. The estimated scrap loss of foil for this process is about 8%. The estimated resources to meet these needs are the following:

Electrode Slitting	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
	8,170,000	4 per shift	2.0 mil\$ total	300
	m ² /y cell area			

*Total cost including installation

5.3.7 Final Electrode Drying

In the absence of electrolyte, no harm is done by exposing the electrodes to normal factory air; however, the electrodes must be dried by heating under vacuum prior to cell assembly. Maintaining extremely low moisture conditions during cell assembly is believed to be very important in achieving long battery life. The final drying step coupled with dry room conditions ensures minimal moisture content in the final product. The pertinent processing rate in determining the resources necessary for drying is the total amount of active materials processed per year (other electrode materials are approximately proportional), which for the baseline plant is 2,950,000 kg/y or 3,275 kg/shift. The individual electrodes exiting from the electrode slitting process are separated into stacks by polarity, loaded into vacuum drying ovens, dried for several hours, and unloaded directly into the dry room. The estimated resources to meet these needs are the following:

Electrode Drying	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
	2,950,000 kg/y	2 per shift	1.6 mil\$ total	300
	active material			
Dryer capacity			600 kg/shift	
Number of dryers			8	
Maximum rate			4,320,000 kg/y	

*Total cost including installation

5.3.8 Control Laboratory

The purpose of the control laboratory is to ensure that the raw materials and the electrodes being fabricated meet specifications. Laboratory personnel collect or supervise collection of samples and carry out analyses. The estimated resources to meet these needs are the following:

	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
Control Lab	869,000 kWh/y	4 per shift	1.5 mil\$ total	300

*Total cost including installation

5.3.9 Cell Stacking

The cells are assembled in four steps, which are carried out in a dry room. The first of these steps is cell stacking. The primary rate factor that determines the cost for all steps in cell assembly is the number of cells assembled per year. For cell stacking an additional cost factor is the capacity of the cells; large cells usually require more electrodes of larger area and thus a more capable, faster cell stacking machine. The method used to determine the extra costs of stacking equipment is detailed in Table 5.4. The capacity of the cells is deemed to have only a minor effect on the other steps in cell assembly and is not taken into account for those steps. The electrodes are inserted in a folded separator sheet, the positive electrodes tabs protrude on one side and the negative electrodes tabs on the other. As in other parts of the plant, excess capacity is provided to allow catching up after unscheduled downtime. The estimated resources to meet these needs for the baseline plant are the following:

Cell Stacking	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
	6,320,000 cells/y	5 per shift	4.0 mil\$ total	600
Stacking rate			5 cells/min	
Number of units			4	
Maximum rate			8,640,000 cells/y	

*Total cost including installation

5.3.10 Current Collector Welding

The current collector tabs for the negative and positive electrodes are welded to their respective terminals by ultrasonic welding. This procedure achieves a connection of near-zero resistance and avoids overheating the electrodes during the welding process. The estimated resources to meet these needs are the following:

Tab Welding	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
	6,320,000 cells/y	5 per shift	4.0 mil\$ total	600
Cell rate			5 cells/min	
Number of units			4	
Maximum rate			8,640,000 cells/y	

*Total cost including installation

5.3.11 Enclosing Cell in Container

The aluminum foil layer in the pouch container is sufficiently thick (100 microns default thickness) to permit the use of stiff, pre-shaped pouch halves. The pouches are assumed to be purchased as finished parts. Each cell is enclosed in these containers, which are then partially

sealed prior to injecting electrolyte. The estimated resources to meet these needs are the following:

Enclosing cells	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
	6,320,000 cells/y	3 per shift	3.0 mil\$ total	600
Cell rate			5 cells/min	
Number of units			4	
Maximum rate			8,640,000 cells/y	

*Total cost including installation

5.3.12 Electrolyte Filling and Cell Sealing

At this station, the cells are evacuated, filled with electrolyte and temporarily sealed. The estimated resources to meet these needs are the following:

Filling & 1 st Seal	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
	6,320,000 cells/y	5 per shift	5.0 mil\$ total	900
Cell rate			5 cells/min	
Number of units			4	
Maximum rate			8,640,000 cells/y	

*Total cost including installation

5.3.13 Dry Room Management

Excellent dry-room atmosphere is required for lithium-ion cell assembly. A maximum dew point temperature of -40 °C is maintained in the room. The load on the dry-room drying apparatus is determined by diffusion of water vapor through the walls, entry of air through the air locks, the number of workers in the room, and the need to admit some fresh air to limit the build up of contaminants such as electrolyte solvent vapor. These load factors are approximately a function of the room area. Because of the importance of the proper functioning of the dry room, two workers are on duty at all times to monitor its performance. The equipment for circulation and purification of the dry air will be located outside of the plant building, adjacent to the dry room. The estimated resources to meet these needs are the following:

	Operating Area	Direct Labor	Capital Equip.*	Air Locks, m ²
Dry Room	$3,000 \text{ m}^2$	2 per shift	20.0 mil\$ total	100

*Total cost including installation

5.3.14 Formation Cycling, Final Cell Sealing and Charge Retention Testing

Formation cycling is expensive because it takes considerable time and each cell must be monitored separately. For plants to be operated in 2020, we expect some improvements from present day operations because of the urgency to improve and thus save cost. We project that the entire formation cycling and testing can be done in two shifts. These operations consist of charging the cell, discharging to full depth to measure capacity and impedance, followed by fully recharging the cells. These tests will be carried out in large temperature controlled cycling units that test 500 cells simultaneously, monitor each cell and automatically identify failed cells. The capital cost of the cycling equipment is primarily a function of the annual number of cells to be tested, but to a lesser extent on the capacity of the cells.

The short-term testing described above does not detect cells that have self-discharge rates that are slightly above normal, which could lead to catastrophic failures later. To detect such defects, the cell charge is topped off and the cells are stored for two weeks and then checked for loss of charge. Most of the test period is spent in large racks in compact arrays, without electronic monitoring. Incidentally, the two-week long self-discharge testing requires less floor space than for formation cycling, which lasts only two shifts.

The final cell sealing occurs between the formation cycling and charge-retention storage test. Gas generated during formation cycling may accumulate in the reservoir space that was created during the temporary sealing step. This gas is removed by creating the final seal below the reservoir and trimming off the unwanted portion.

	Rate Factor	Direct	Capital Equip.*	Plant Area,
		Labor		m^2
Formation Cycling	6,320,000 cells/y	8 per shift	30.0 mil\$ total	2200
Cell capacity			40 Ah	
Number of cyclers			35	
Cells per cycler			500	
Length of test			2 shifts	
Testing capacity			7,875,000 cells/y	
Final Cell Sealing	6,320,000 cells/y	2 per shift	2.0 mil\$ total	450
		_		
Charge Retention	6,320,000 cells/y	3 per shift	4.75 mil\$ total	900
Testing rack capacity			500 cells	
Racks per stack			5	
Number of racks			750	
Length of test			14 days	
Testing capacity			8,040,000	

The estimated resources to meet these needs are the following:

*Total cost including installation

5.3.15 Module and Battery Assembly

Approximately 5% of the cells are expected to fail the formation cycling and charge-retention tests and these are sent to the Rejected Cell and Scrap Recycle section. The accepted cells (6,000,000 finished cells per year) are assembled into modules by attaching the terminals through laser welding or mechanical joining with spring loaded devices. Electronic circuit packs are attached that occupy about the same volume as a cell. An aluminum heat conductor is placed around every cell. These operations are carried out at four automated stations each capable of

handling about 280 cells per hour. For the module design being cost estimated in this model, the module is enclosed in an air-tight aluminum container by double seaming. The processing rate that determines the cost of module assembly is the number of finished cells that must be handled per year.

The finished modules are assembled into battery packs with the aid of automated stations. The total cost of these stations is dependent mainly on the number of battery packs to be assembled per year (100,000 for the baseline plant), but to a lesser extent on the number of modules per pack. After assembly, the packs are moved to testing stations where they are discharged as a final check of impedance and to lower the state of charge to a level suitable for shipping. The estimated resources to meet these needs are the following:

	Rate Factor	Direct	Capital Equip.*	Plant Area,
		Labor		m^2
Module Assembly	6,000,000 cells/y	6 per shift	6.0 mil\$ total	600
Number of stations			4	
Cells/h/station			280	
Capacity			8,060,000 cells/y	
Battery Pack Assembly	100,000 packs/y	3 per shift	3.0 mil\$ total	450
Modules/pack			4	
Number of stations			3	
Packs/h/station			6	
Capacity			130,000 packs	
Battery Pack Testing	100,000 packs/y	3 per shift	3.0 mil\$ total	450
		_		

5.3.16 Rejected Cell and Scrap Recycle

Scrap is generated in preparing the electrodes and by the rejection of 5% of the cells that go through formation cycling and charge-retention tests. This scrap is gathered and packaged for shipment for recycling of the materials having value. No credit is taken for the value of the scrap in this model except that the costs of gathering, sorting, packaging and shipping are understated by about that value. The main factor in determining the cost of scrap recycle is the number of cells rejected, which have to be disassembled to recover the scrap, a labor intensive process. The yields of materials in the various processing steps are shown in Table 5.5.

Material	Material	Coating	Electrode	Cell	Electrolyte	Total
	Mixing		Slitting	Stacking	Filling	
Positive Electrode	99	95	99	99		92.2
Negative Electrode	99	95	99	99		92.2
Positive Current Coll.		99	92	99		90.2
Negative Current Coll.		99	92	99		90.2
Separator				98		98.0
Electrolyte					94	94.0

Table 5.5 Materials yields during electrode and cell fabrication

The estimated resources needed for scrap recycle are the following:

	Rate Factor	Direct Labor	Capital Equip.*	Plant Area, m ²
Scrap Recycle	6,320,000 cells/y	5 per shift	2.5 mil\$ total	600
Scrap rate			441 kg/shift	

5.3.17 Baseline Plant Summary

The processing rates and the primary cost factors for the baseline plant are summarized in Table 5.4. The main processing rate for each step is shown in the second column. The requirements for direct labor, capital equipment and plant area, which are shown in detail in the subsections above, are summarized in the table. It is seen that the plant requires a total of 90 workers per shift, \$127,450,000 worth of capital equipment, and 15,425 square meters of plant area to manufacture the baseline battery at a rate of 100,000 battery packs per year.

5.4 Adjustment of Costs for Varying Production Volumes

Production volume may affect the end price of the battery in two distinct ways. First, the user of the model may change the annual production volume and every processing step will be affected. Somewhat differently, as the performance requirement and thus design is changed, the production of individual steps will change in non-uniform ways. As noted in Table 5.4, there are many processing rates that must be considered in addition to the overall number of battery packs manufactured per year. Each of these rates affects the costs of one or more steps in the process and may have no effect upon the costs of other steps in the process. For instance, when the user of the model increases the power of the battery packs without increasing the number of cells or their capacity, the model increases the area of the cells and decreases the electrode coating thicknesses. Such changes would result in an increase in the cost of the process. It would have no effect on the cost of mixing the materials to be coated because the amounts of these materials per battery back are unchanged under the assumed conditions.

The general approach to cost estimation of multiplying a known cost by the ratio of processing rates raised to a power has also been applied to the capital cost of individual items of equipment.⁴⁹

$$C = C_0 (R/R_0)^p \tag{5.2}$$

Here, C_0 is the capital cost of an installed equipment item designed for the baseline processing rate, R_0 . The power factor, p, relates the capital investment cost and the processing rate for the manufacturing step.

If the value of p were 1.0, it would imply that the cost of the equipment item, or the equipment items if there are several in parallel, would be directly proportional to the processing rate. However, the value of p for the cost of equipment is frequently about 0.6 to 0.7 for many

manufacturing process steps because the equipment is larger for the higher processing rates and its cost is less than if it were directly proportional to the processing rate. For process steps requiring the addition of many identical pieces of equipment for scale up, such as may be true for formation cycling of battery cells, the value of p may be as high as 0.9. The value of p is unlikely to reach 1.0 because the equipment cost includes installation, for which there is some savings even in installing multiple units of the same processing capacity. The relationships between cost and processing rate for two-fold and three-fold rate changes are illustrated in Table 5.6.

$C/C_o = (R/R_o)^p$				
	Cost Ratio, C/C_{o}			
Scale Factor, p	$R/R_{\rm o}=2$	$R/R_{\rm o}=3$		
0.25	1.19	1.32		
0.3	1.23	1.39		
0.4	1.32	1.55		
0.5	1.41	1.73		
0.6	1.52	1.93		
0.7	1.62	2.16		
0.8	1.74	2.41		
0.95	1.93	2.84		
1.0	2.00	3.00		

Table 5.6 The effect of processing rate (R) on cost for various scale factors

Similar equations have been applied for determining the effect of processing rate on the annual hours of labor and the plant area required for a manufacturing step. In general, the value of p is low for the labor equation, usually only 0.4 to 0.5, because only a relatively small addition to the labor crew permits operation of larger equipment or of operating several more units of the same processing capacity.²⁴ The value of p for the plant area required for a processing step is slightly less than that for equipment. The floor area required for larger equipment or for more equipment items of the same size is proportionately less than the increase in the processing rate because of the more efficient use of the space occupied by the equipment and the savings in aisle area.

The value of the scale factors (i.e. p factors) for labor, capital equipment, and floor area were estimated for each of the processing steps (Table 5.4). The scale factors selected for the direct labor requirement are usually only 0.4 to 0.5, which indicates considerable unit cost reduction for increasing the plant throughput.

For most processing steps, increasing the processing rate beyond that in the baseline plant would result in a decision to increase automation or use faster equipment to mitigate the costs of higher levels of throughput. Decreasing the processing rate would have the opposite effect. Some steps in the process such as cell stacking, welding of current collectors, and formation cycling do not appear to be easily automated beyond the level intended in the baseline plant and, thus require a higher value for the scale factor of 0.8. This higher scale factor results in achieving fewer reductions in the cost per battery pack with increasing production volume. Additionally, a higher

p factor results in a less severe penalty for lower production scale for an individual step in the process.

There are five steps for which the cost of the capital equipment is affected by factors other than the main processing rate for the process step. These are discussed in the footnotes at the bottom of Table 5.4. For these steps, the costs that have been adjusted for the changes in the processing rate from the baseline rate are further adjusted to take into account the other cost factors. The cost of the coating equipment is adjusted for the amount of solvent to be driven off the positive and negative electrodes; thicker coatings need longer, more expensive ovens to drive off the additional binder solvent or the coater most be operated at lower speeds. The costs of the cell stacking equipment and the formation cycling equipment, for which the main cost factor in both cases is the number of cells to be fabricated annually, are also adjusted for the capital equipment for battery assembly is primarily a function of the number of cells in the battery, but it is also a function of the number of modules that must be interconnected. This dependence is accounted for in the model with an additional multiplying factor.

A breakdown of the baseline plant capital equipment costs listed in Table 5.4 is illustrated in Fig. 5.3. The largest costs for capital equipment are for formation cycling and testing, cell assembly (in the dry room) and electrode coating. These capital costs are likely to be dominant in any lithium-ion battery plant in the near future.



Figure 5.3 Breakdown of installed capital equipment costs for the baseline plant

5.5 Plant Investment Costs

In this model, the calculated investment costs are defined as those directly related with building and operating the plant (Table 5.7). Other costs that may require investment, such as research and development, are added separately to the unit cost of the battery. The largest investment cost is for the installed capital equipment. Each cost item for the battery under design is adjusted from the estimate of the baseline plant. The plant cost is done in a similar way with a cost of \$3,000 per square meter (\$280/sq. ft) including land and utilities. The high cost for land and utilities accounts for both the area of the manufacturing facility as well as other land requirements such as office buildings and waste water treatment requirements. Launch costs include plant start-up, employee training and materials that are lost or recycled in early stages of production, beyond the normal amounts. Launch costs are estimated to be 5 % of annual materials costs plus 10 % of annual direct labor and variable overhead (Section 5.6). Working capital is needed to cover the costs of payroll, receivables, and the inventories of raw materials, work in progress and finished product. These working capital costs are partially offset by bills that are payable. We estimate the working capital to be 15 % of the annual variable costs.

Investment Costs	Description	Method of Calculation
Capital Equipment	Equipment costs including	Estimates of costs for each
	installation	processing step at baseline rates
		adjusted for actual rates.
Plant Floor Space	Space includes aisles and space	Estimates of costs for each
	for unfinished processing	processing step at baseline rates
	inventory plus land and utility	adjusted for actual rates.
	costs.	
Launch Costs	Plant start-up, training, out-of-	5% of annual materials cost,
	spec product.	10% of direct labor plus
		variable overhead.
Working Capital	Cash to meet payroll,	15% of annual variable costs.
	receivables, inventories of raw	
	materials and of unfinished and	
	finished product, minus	
	payables.	

Table 5.7 Battery pack manufacturing investment costs

5.6 Unit Costs for Battery Pack

The unit costs of the battery pack are calculated as summarized in Table 5.8.

5.6.1 Variable Costs

The costs of the materials and purchased items are based on the costs discussed in section 5.2, and the annual amounts of materials are adjusted for the yields of materials (section 5.3) and yield of cells. The direct labor is the sum of the labor cost for each step in the process, which are

each calculated for the baseline plant and adjusted for the rate associated with the battery under study. Variable overhead is the cost of indirect materials and labor, utilities, and plant maintenance. It is estimated to cost 40 % of direct labor costs and 20 % of total depreciation.

5.6.2 Fixed Expenses

Fixed expenses include General, Sales, and Administration (GSA), research and development, and depreciation. The cost of GSA includes the plant office, taxes on income and property, cost of sales and insurance. It is estimated by the model as 25 % of direct overhead and depreciation. Research and development (R&D) must be carried out to ensure that the battery packs that are produced in the plant and the means of production continue to be competitive in the world market with respect to performance and price. The greater the investment in the plant and its equipment, the greater is the need to be successful in the R&D effort. Thus, the expenditure has been set at 40 % of the depreciation expense. Depreciation expense provides funding available for future investment in this plant or another venture to replace deteriorating plant and equipment. The equipment and plant are depreciated at straight-line rates for 6-year life (16.7 % per year) and 20-year life (5 % per year).

Variable Costs	Description	Method of Calculation	
Materials and Purchased	All materials and purchased	Based on prices of materials,	
Items	items in finished product and	cost equations for purchased	
	lost in processing.	items and yields.	
Direct Labor	Labor costs for operations and	Estimates of costs for each	
	immediate supervision.	processing step at baseline rates	
	_	adjusted for actual rates.	
Variable Overhead	Indirect materials, labor,	40% of direct labor cost plus	
	utilities, plant maintenance	20% of depreciation	
Fixed Expenses			
General, Sales, and	Plant office, taxes on income	25% of direct labor and variable	
Administration (GSA)	and property, cost of sales and	overhead plus 25% of	
	insurance expenses.	depreciation.	
Research and Development	On-going research needed to	40% of depreciation	
	upgrade product and maintain		
	competitive position.		
Depreciation	Provides funds for new	16.7% of capital equipment cost	
	investments to replace those in	plus 5% of plant floor space	
	current equipment and plant.	cost.	
Profit	Return on invested capital after	5% of total investment costs.	
	taxes.		
Warranty	Funds set aside for reimbursing	5.6% added to price based on	
	customers for battery pack	present worth of projected	
	failures.	payments.	

 Table 5.8 Unit cost of battery pack

5.6.3 Profits

The profit goal for this type of venture varies with the financial structure of the company, especially regarding long-term debt. For the model, the profit is set to provide a 5 % return on the total investment, which is an approximate average for mature manufacturing as vehicle battery production is expected to be in 2020. In general, the chosen cost structure and the resulting margin are similar to a Tier 1 supplier in the automotive industry.

5.6.4 Battery Pack Warranty Costs

If a battery module or an entire pack fails, the replacement will cost much more than the original price paid by the OEM. It is important that such events are rare, but provision must be made to reimburse the vehicle owner, especially in the early years of the projected battery life. The extra costs of replacing the battery will result from labor for testing and replacing the battery, inventory costs for stocking replacement batteries, and servicing the battery controller if the new battery is slightly different than the old battery. It is likely that the battery manufacturer will be responsible for the cost of the new battery, which we assume will be equal to the cost of the original battery. The other costs of replacing the battery, to the extent that they are covered by the warranty, are assumed here to be covered by the automobile manufacturer and the dealer. The goal for average battery life is 15 years and a warranted life of 10 years, with full replacement in the first five years and shared cost of replacement for the last five years seems appropriate. The vehicle owner would pay an increasing share of the cost from between 0 % at 5 years to 100 % at 10 or more years. With these assumptions, the cost to the battery manufacturer will be equal to the present worth of the future costs of the new battery or modules as provided in the warranty. The rate of battery failure will vary over the life of the battery with a slightly higher rate early in life, then a low failure rate followed by a gradually increasing failure rate. For purposes of calculation we assume a failure rate of 1.0 % per year throughout the warranty period. With an internal rate of return of 8 % and calculated on a monthly basis, the present value of the future costs would be about 5.6 % of the price of the battery before adding the warranty cost.

5.7 Summary of Baseline Battery Cost

The spreadsheet version of the model, which is discussed in more detail in sections 6 and 7, provides a summary sheet which is illustrated in Table 5.9 for the cost of the baseline battery and that of three others. This breakdown of the battery costs, with a brief summary of the design values, illustrates the effects of the cost factors. The second battery has twice the power of the baseline battery and the third battery has the same power as the baseline battery, but twice the capacity. The number of cells is the same for each battery. The energy storage is slightly higher for the battery with double power because the voltage would be slightly higher during the discharge to determine capacity. The battery with double the capacity has about the same number of electrodes; although, the electrodes are longer and wider in dimension, because the cell thickness is maintained. This results in a slightly higher resistance in the current-collector structure. The higher impedance lowers the voltage during the discharge capacity measurements and results in slightly less than twice the energy storage of the baseline battery.

Calculated Batte	ery Parameters		Baseline	Double Power	Double Capacity	Double Modules
Vehicle electric range	e, miles		24.4	24.6	48.5	48.6
Number of battery pa	cks		1	1	1	1
Packs in series of	r parallel					
Number of cells per p	ack		60	60	60	120
Battery system total	energy storage, kV	/h	8.7	8.8	17.3	17.3
Cell capacity, Ah			40.0	40.0	80.0	40.0
Cell group capacity, A	Ah 🛛		40.0	40.0	80.0	40.0
Module capacity, Ah			40.0	40.0	80.0	40.0
Pack capacity, Ah			40.0	40.0	80.0	40.0
Battery system capa	city, Ah		40.0	40.0	80.0	40.0
Nominal battery syste	em voltage (OCV at	50% SOC)	221	221	221	442
Battery system powe	r at target % OCV,	kW	50.0	100.0	81.0	85.9
Required battery syst	em power, kW		50.0	100.0	50.0	50.0
Target % OCV at full	power		80.0	80.0	80.0	80.0
% OCV at full power a	adjusted for thickne	ess limit	80.0	80.0	88.9	89.6
Battery system volum	ne (all packs), L		37.4	43.2	61.3	66.2
Battery system mass	s (all packs), kg		63.3	78.1	112.3	117.7
Cooling system powe	er requirement, W		616	251	348	335
Investment Cost	ts					
Capital equipment co	st including installa	tion. mil\$	128	146	158	205
Building, Land and Ut	ilities					
Area m ²			15 478	18 401	19 204	23 791
Cost $\frac{\varphi}{m^2}$			2 000	2 000	3,000	2 000
Building investme	ot mil¢		3,000	55.2	57.6	5,000 71 /
Building investment	π, ππφ		40.4	55.2	57.0	71.4
Pate: 5% of direct	onnual matariale	10% of other	annual costs			
Total million				13.02	16.46	18.00
Working capital (15%	of annual variable	costs) mil\$	27.71	35.52	10.40	10.05
Total investment mil		οστο), πηφ	212.11	2/19/15	277.07	344.20
Summony of Uni	t Casta ¢		212.10	243.43	211.51	544.20
Summary of Om	ι σοδιδ, φ		4.045	4 700	0.000	0.040
Materials			1,245	1,706	2,329	2,342
Purchased Items			397	426	481	6/1
Direct Labor			113	130	130	162
Variable Overnead			92	106	111	140
General, Sales, Admi	nistration		110	127	133	170
Research and Develo	pment		94	108	117	151
Depreciation			236	270	292	377
Profit			106	125	139	172
vvarranty			134	168	209	234
Price to OEM for battery pack, \$		2,528	3,166	3,941	4,421	
Pack integration (BIVI	S & Disconnects),	\$/раск	395	395	395	475
Total battery cost to C		ι. Φ	2,923	3,561	4,336	4,896
Additions to AC for thermal management, \$		240	200	200	200	
Iotal cost to OEM for complete system, \$		3,163	3,761	4,536	5,096	
Price to UEM for modules for one pack, \$			2,364	2,977	3,734	4,204
Error bars on price to		аск, %				
Potential positive	error, %		21	10	31	26
Potential negative	error, %		10	10	16	15

Table 5.9. Summary of results for cost of baseline battery and that of similar batteries with double the power and double the capacity of the baseline battery

Doubling the power does not add as much cost to the materials and purchased parts as doubling the cell capacity. Most of the labor costs for the first three batteries are similar with the major difference being for the labor cost for electrode processing. The double power battery requires greater labor costs principally for coating the larger electrode area. Capital equipment and depreciation costs are higher for both the high power and high capacity battery packs. The increases in capital equipment cost for the high-power battery are for coating, calendering, materials handling and vacuum drying equipment. For the high-capacity battery, the main additional capital equipment costs are for the materials mixing, binder solvent recovery, cell stacking and formation cycling steps in the process.

Overall, doubling the power of the battery increases the price of the battery pack by only 25 %. Doubling the capacity of the cells increases the cost by 56 %, considerably more than for doubling the power. Alternatively, doubling the number of baseline cells and modules within a larger battery jacket (two rows of modules instead of one, twice the voltage, energy, and power) would increase the cost by 75 %.

The summary of unit costs for the baseline battery pack, which is shown at the bottom of Table 5.9, is illustrated in Fig. 5.4. The materials and purchased items are the largest costs for the battery. For larger levels of production, these costs are even more dominant because the scale factors for these items are close to one.



Figure 5.4 Breakdown of unit costs for baseline battery with total price to OEM of \$2528. The total battery cost to the OEM, including pack integration components but excluding thermal management external to the pack, is \$2,923.

5.8 Uncertainties in Price Estimates

The potential uncertainty in estimating the price of future Li-ion batteries depends upon many factors. In BatPaC, the error bars for the 95% confidence intervals for the price charged to an OEM for the battery pack are automatically calculated. The uncertainty ranges are specific for the performance requirements of the battery. The three major categories of uncertainty in the default calculation of battery price are:

- Input costs of materials and capital equipment
- The maximum allowable thicknesses of the electrode coatings
- The maximum capacity of single cells

An in-depth variation study analyzing the contribution of these categories to the battery price to the OEM has been completed. The details and methodology used in that study may be found in the Appendix.

5.8.1 Materials and Capital Equipment

A variation study was made of the cost inputs for the top eight contributors to total battery price including the active materials, copper current-collector foil, electrolyte, separator, and SOC controllers (see Appendix). The costs of capital for electrode coating and formation cycling were also varied. Both of these are subject to considerable variation due to unforeseen quality difficulties or advances in process throughput. The high, median, and low costs were considered for each of the factors, sometimes over 100% higher in value. Normal or log-normal random distributions of these values were used to calculate a population of batteries with differing price. Analyzing the resulting uncertainty in end battery price, a range of +/- 10% was found to capture the 95% confidence interval of the total price for several different cell chemistries and for HEV, PHEV, and EV battery packs.

5.8.2 Electrode Thickness

Most of the current transportation Li-ion batteries have electrode coatings of about 20-60 micron thickness; whereas, some consumer electronic applications may have 120-micron thick electrodes. As batteries are sought for vehicles with useful electric ranges (low power-to-energy requirement), there is a strong incentive to save cost by increasing the thickness of the electrode coatings as discussed previously in section 3.6.1. In BatPaC, a 100-micron default electrode thickness limitation is assumed. If the calculated optimal thickness is less than 100 microns, then the optimal calculated thickness is used. If the optimal thickness is greater than 100 microns, BatPaC increases the area of the cell to meet the energy requirement while maintaining an electrode thickness of 100 microns. After continued engineering development, the appropriate thickness limitation may be greater or less than the default assumption. The model estimates this uncertainty by calculating the total costs for batteries having electrode thicknesses limits at the 95% confidence limits, namely 50 and 150 microns for PHEV cells and 70 and 200 microns for EV cells. EV cells typically operate at lower P/E ratios than PHEVs and thus are more likely to successfully utilize electrodes of larger thicknesses. The costs so calculated determine negative and positive percentage uncertainty added to the base value of 10 % discussed in section 5.7.1.

5.8.3 Cell Capacity

Vehicle designs requiring batteries with large energy storage (e.g. EVs) commonly rely on high capacity cells or cell groups, exceeding 50 Ah. Cells of this size and larger (e.g. 200 Ah) exist today for some applications and cell chemistries. However, an OEM may choose to use smaller cells grouped by parallel connection. These smaller cells may have already been demonstrated to meet the performance, life, and safety requirements for a particular application and thus are a more attractive choice for the OEM. It is difficult at this time to predict the largest cells that will be practical for a particular application in the near future and, thus, BatPaC does not limit the cell capacity. However, as the cell capacity calculated to meet the energy requirement increases, the likelihood of the designers using parallel connected cells in a group also increases. This uncertainty in additional cost is taken into account by increasing the size of the positive error bar. Additional parallel cells increase the cost because of the increased number of cell terminals, interconnects, and formation cycling units in the manufacturing facility. Details on these costs are discussed in section 7.3.

To calculate the cost uncertainty from lowering the capacity of individual cells, BatPaC sets a capacity of 50 Ah below which there is no addition to the error bar. At 100 Ah or above the error bar is that for the additional cost of using half the cell capacity and doubling the number of cells with two cells in parallel. Between 50 and 100 Ah the error bar is calculated as 2% of the additional cost for the smaller capacity cells in parallel for each Ah above 50 Ah.

5.8.4 Example of Contribution to Calculated Uncertainty

The uncertainties in the cost estimates are calculated in the worksheet "Error Bars", in which, BatPaC recalculates the cost of the battery pack with the maximum electrode thicknesses and the capacity set at what we estimate to be the 95% limits and compares the results with that for the default values, resulting in contributions to both positive and negative error bars. An example is illustrated in Fig. 5.4 for a 120 kW 360V EV battery based on the NMC441-Gr chemistry having a vehicle range of 80 to 200 miles. As the range is increased, both the thickness of the electrodes and the capacity of the cells increase. No uncertainty from the electrode thickness limitation is added for calculated electrode thickness less than 70-micron. As the thickness increases, the positive uncertainty increases. Above 100 microns in thickness (default assumption), a negative uncertainty is added on the assumption that the actual thickness limit may be as high as 200 microns. Similarly, the effect of the uncertainty derived from the cell capacity is not a factor for capacities less than 50 Ah. A contribution to the positive uncertainty is accounted for as the capacity increases above that level. In summary, for this example, the battery price uncertainty ranges are plus 10% and minus 10% for the vehicle with 80-mile range, and plus 26% and minus 18% for the vehicle with 200-mile range.



Figure 5.5 Effects of electric-vehicle range on price and error bars for battery pack with NMC441–Gr cell chemistry, 120-kW 360 V.

5.9 Breakdown of Costs

BatPaC assists in showing the relative importance of cost items by providing cost breakdowns from several perspectives as detailed in the Cost Breakdown worksheet. The costs are distributed in several ways and then graphed with pie charts to illustrate the relative importance of the cost items. This is a useful way of judging the importance of cost items.

This worksheet is illustrated below with the sample cases provided in BatPaC v2.0 as distributed. The sample batteries are PHEV batteries all with 60 LMO-G cells and power of 60 kW. They range in energy from 4 to 16 kWh and provide an electric range of about 10 to 42 miles for vehicles requiring energy of 250 Wh/mile.

The first set of graphs for Battery 1 (4.0 kWh) and Battery 7 (16.0 kWh) shows the overall distribution of costs as calculated in the "Summary of Results" worksheet. (The modeler may change the selection of battery on the chart by selecting the pie image on the chart and then changing the Excel column for the cost values.)



Figure 5.6 Overall distribution of costs of manufacturing battery packs, not including pack integration system.

These graphs illustrate that materials and purchased items compose a large fraction of the total battery pack price for a mature battery plant producing 100,000 packs per year. This effect is stronger for the 16-kWh battery (Battery 7), which has a large fraction of expensive cell materials, than for the smaller 4-kWh battery.

The second set of graphs illustrates the breakdown of costs for the materials and purchased items only. For the 16-kWh battery (Battery 7), the costs of active materials and electrolyte are substantial. It should also be noted that the cost of separator material is greater than the total costs of the current collection foils. The fraction of cost required for active materials is larger for the 16-kWh battery than for the 4-kWh battery because the cell capacity is larger and the electrodes are thicker.



Figure 5.7 Distribution of materials and purchased items for the battery pack.

Sections 5.5 and 5.6 discuss the derivation of fixed and variable overhead costs by the use of multiplying factors on the costs of (1) materials and purchased items, (2) direct labor, (3) capital equipment, and (4) building, land and utilities. These overhead costs can be redistributed back to their origins by applying the constant multipliers shown in Table 5.10 which is a reproduction from the "Cost Input" worksheet.

The next set of charts (Fig 5.7) in the printout of the Cost Breakdown worksheet, shows the distribution of the price of the battery pack to materials, purchased items, and the individual manufacturing processes and operations.

 Table 5.10 Table for converting overhead costs back to their origins

Multipliers for Overhead to Basic Costs					
Basic Cost Factors	Multiplier				
Materials and purchased items, \$/pack	1.0666				
Direct labor, \$/pack	1.8665				
Capital equipment (100K packs/yr*)	3.8764				
Building, land, utilities (100K packs/yr*)	1.5325				
*For other production rates multiply by 100,000/rate					



Figure 5.8 Distribution of overhead costs to materials, purchased items, and individual processes.

The materials and purchased item costs have a small effect on overhead cost, namely launch costs and working capital costs. Thus we have distributed back to the materials and purchased items a small amount, which accounts for the fraction of the cost of materials and purchased items being slightly higher in the charts of Fig. 5.7 than in those of Fig. 5.5. The other costs are redistributed to the individual processes and operations.

The costs of the three major manufacturing operations of (1) electrode processing, (2) cell assembly, and (3) formation cycling, testing and sealing estimated by BatPaC for a

manufacturing scale of 100,000 battery packs per year in 2020 is considerably lower than their actual costs at the time this report is in writing. If these projected costs are reached, there can be only small percentage reductions in cost after that by further optimization and automation for even larger scales of production, unless there are substantial reductions in materials costs.



In Fig. 5.8, the costs are distributed to the basic cost factors of (1) materials, (2) purchased items, (3) labor, and (4) capital equipment and building by means of the factors in Table 5.10.

Figure 5.9 Redistribution of costs to basic cost factors

The total effects of the direct labor costs are greater than indicated in Fig. 5.5 because of the influence of labor costs on overhead as shown in Fig. 5.8. Foreign battery plants may have lower labor cost than plants in the United States, which may give them an advantage even considering shipping costs. However, the advantage would be small and further automation may overcome such an advantage.

Clearly, the major opportunities for cost reduction are in the area of materials costs. Improvements may be in lower costs per unit weight of material or in higher performance, greater power or greater capacity per unit weight for materials of nearly the same cost as set in BatPaC.
6 Description of the Spreadsheet Model and Instructions for Use

6.1 Background

Historically, the model has been based on Microsoft[®] Office Excel spreadsheets. The flexibility afforded by a spreadsheet approach has been extremely useful to the development of the calculations. Until recently, the model had been in a constant state of development. Changes to parameters and equations were made rapidly and frequently. The publication of the BatPaC v1.0 report represented the first time a version of the model was be "frozen" for open distribution to the public. Advances will continue to be made with the model beyond this current v2.1; however, distributions of the revised model will be made in an orderly fashion rather than the continuous improvement approach taken over the last number of years.

6.2 Instructions

The following subsections are a brief explanation of how one may operate the spreadsheet based model. The user is advised to save the original document separately as a back-up copy. Corruption of the calculation is possible and will likely occur during use by someone unfamiliar with the model.

6.2.1 Enabling Calculation

This Microsoft[®] Office Excel workbook requires the use of iteration. The iterative function may be enabled by going to File > Options > Formulas. Check the box next to "Iteration" and change the maximum number of iterations to 1000 (Figure 6.1). Perhaps most importantly, ensure the calculation is set to automatic and not manual. If the iteration is not turned on, the software will present an error complaining about circular references. If the model is opened while a different Excel spreadsheet is in use, the software will also warn of an error. Simply close all Excel windows except for the model; alternatively, one could re-enable the iterative function as discussed above.

6.2.2 Chem Worksheet

The cell chemistry is selected by copying the system designated at the top of a column, for instance NCA-G in cell F4, pasting it into cell E4 (Figure 6.2). Any of the values in column E can be overridden by entering the desired value in column M. For example, the maximum electrode thickness may be overridden by placing a new value in cell M53. The selection of the cell chemistry also includes the associated prices at the bottom of the page. These prices can also be overridden by entering the desired values in column M. A full screen shot of the system selection worksheet is in Figure 6.3.

6.2.3 Battery Design Worksheet

The Battery Design worksheet designs seven or more batteries for any type of electric-drive vehicle (Figure 6.4–6.6). The calculated designs are specific for the end battery requirements specified by the user. From the result, the amounts of materials and the purchased items required for manufacture are easily available to be used in the manufacturing cost calculations found on

subsequent worksheets. Although a cell and module format is assumed, the exact format (prismatic, pouch, can, etc) of the battery does not have a dominant effect on the cost for a set cell chemistry system. Our experience teaches us that the amounts of electrode materials and the number, capacity and electrode area of the cells, are the determining cost factors. Nevertheless, a specific design format was selected and is shown on the Cell Design worksheet to provide a basis for calculating the entire cell and battery related costs.



Figure 6.1 Automatic iteration must be enabled for the spreadsheet model to function.

				BatPa	ac vz.r - Mici	osoft Excel								-
Home Insert Page Layout	Formulas Data	Review	View										V	() -
	D F	F	G	н		. I	К		М	N	0	P	0	R
, B 0	0	C	ell Chemis	strv			i.	-					~	
	Selected			,										
	System			Default	Values				Override					
	L MO-G	NCA-G	NMC441-G	NMC333-G	I EP-C		LMO-G	Other	Valuee					
locitivo Electrodo	LM0-0		111110441-0	NWC000-G	611-0	LINO-LIO	LINO-0	outer	Valuea					
ositive Electrode	100	160	475	150	150	109	100							
loight %	100	100	175	150	150	100	100							
tive meterial	20	20	20	20	90	90	00							
arbon	6	6	6	6	6	6	6							
inder	5	5	5	5	5	5	5							
inder solvent	NMP	NMP	NMP	NMP	NMP	NMP	NMP							
oid Vol% %	32	32	32	32	50	32	32							
lensity a/cm ³														
ctive material	4 23	4 78	4.65	4.65	3.45	4 23	4 23							
arbon	1.825	1.825	1.825	1.825	1 825	1 825	1.825							
inder	1.77	1.77	1.77	1.77	1.77	1.77	1.77							
legative Electrode														
I/P capacity ratio after formation	12	1.25	1.25	1.25	12	1 10	12							
ctive material canacity mAh/a:	330	330	330	330	330	170	330							
Veight %	000			000	000									
ctive material	95	95	95	95	95	89	95							
arbon	0	0	0	0	0	6	0							
linder	5	5	5	5	5	5	5							
Binder solvent	Water	Water	Water	Water	Water	Water	Water							
oid, Vol% %	34	34	34	34	34	40	34							
Density, g/cm ³														
ctive material	2.24	2.24	2.24	2.24	2.24	3.40	2.24							
Carbon	1.95	1.95	1.95	1.95	1.95	1.95	1.95							
linder	1.1	1.10	1.10	1.10	1.10	1.10	1.10							
Positive Foil														
faterial	Aluminum	Aluminum	Aluminum	Aluminum	Aluminum	Aluminum	Aluminum							
hickness, µm	20	20	20	20	20	20	20							
legative Foil														
laterial	Copper	Copper	Copper	Copper	Copper	Aluminum	Copper							
hickness, µm	12	12	12	12	12	20	12							
N N About PatDaC Cham Patta	n Dacian Cummon	of Booulto		and Taxanta	Manage for advector	Cost Coloriba	Data	to of Modulos	Cash Dave	akdauua	Course Doors	Dbot Cchor	Col	

Figure 6.2 The specific cell chemistry for the battery design is selected on the Chem worksheet.

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A B C D E P 0 H J J L M N 1 Second Second Cell Chemistry Image: Second System Override Override 2 Second Second Second System Default Values Override Override 6 Allos Restricts 80 65 65 6 6 6 6 Second System Override Values 1 Second System 80 65 6		is nome insert Page Layo			REVIEW	view Add-	115						
A B C D E P O H I J K L M N N 2 Selected Selected Selected Overrise Overrise Values Overrise Overrise Values		E4 🔻 (* <i>f</i> x	LMO-G										
Cell Chemistry Vertical Solution of the second of		A B C	D	E	F	G	Н	- I	J	K	L	M	N E
Selected Selected Selected Derivit/Version Override 6 Patter Electrode ILIOS IRCAG INCLAIS INNO233-C EFF-G LIOLTO IRCAG Version 6 Atternetel capecy, mAng 100 175 150 110 100 100 0 7 Disposition 6	1				C	ell Chemi	strv						
Sector Default Values Override 6 Pairter LIGO-G NCA-G NICA-G NICA-G<	2			Selected			, y						
Pasibe Diskup Diskup <thdisk< th=""> <thdisk< th=""> <thdiskup< th=""></thdiskup<></thdisk<></thdisk<>	2			Selected			D-6	1-1				0	
Image: Control of the second	3			System			Default	alues				Override	
B Positive Electrode Post Post Post Post Post Post Post Post	4			LMO-G	NCA-G	NMC441-G	NMC333-G	LFP-G	LMO-LTO	LMO-G	Other	Values	
6 Acte matrix Particle 160 175 150 110 100 100 0 Actes matrix 0 </td <td>5</td> <td>Positive Electrode</td> <td></td>	5	Positive Electrode											
V Magint S. 0 <th< td=""><td>6</td><td>Active material capacity, mAh/g:</td><td></td><td>100</td><td>160</td><td>175</td><td>150</td><td>150</td><td>108</td><td>100</td><td></td><td></td><td></td></th<>	6	Active material capacity, mAh/g:		100	160	175	150	150	108	100			
B Description BB	7	Weight %		0									
9 Cachon 6 <td>8</td> <td>Active material</td> <td></td> <td>89</td> <td>89</td> <td>89</td> <td>89</td> <td>89</td> <td>89</td> <td>89</td> <td></td> <td></td> <td></td>	8	Active material		89	89	89	89	89	89	89			
10 Binder 5 5 5 5 5 5 10 Binder data 32 32 32 32 33 32 32 32 33 32 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 32 33 33 32 33	9	Carbon		6	6	6	6	6	6	6			
II Bucket scheding NMP	10	Binder		5	5	5	5	5	5	5			
12 32 32 32 33 50 32 32 32 12 Market and the material 4.23 4.76 4.85 3.45 4.22 4.22 13 Action material 4.23 4.76 4.85 3.45 4.22 4.22 4.22 14 Action material 1.77	11	Binder solvent		NMP	NMP	NMP	NMP	NMP	NMP	NMP			
13 Details, genth 4.20 4.76 4.65 4.65 3.45 4.23 4.23 16 Carbon 1.025 1.825 1.65 1.65 <	12	Void, Vol% %		32	32	32	32	50	32	32			
14 Acts material 4.22 4.78 4.66 4.65 3.45 4.23 4.23 4.23 15 Brochem 1.27 1.77 <td>13</td> <td>Density, g/cm³</td> <td></td>	13	Density, g/cm ³											
15 Carbon 1825	14	Active material		4.23	4.78	4.65	4.65	3.45	4.23	4.23			
Binder 177<	15	Carbon		1.825	1.825	1.825	1.825	1.825	1.825	1.825			
17 Negative Electrode 12	16	Binder		1.77	1.77	1.77	1.77	1.77	1.77	1.77			
18 MC pacety ratio after formation 12 1.25 1.25 1.2 1.10 1.2 0 Margit S. 330 340 34 34 34 34 34 34 34 34 34 34 34 34 34 34 34 34 34	17	Negative Electrode											
19 Active material capacity, mAvig. 330 330 330 330 330 170 330 21 Active material 95 75 74	18	N/P capacity ratio after formation		1.2	1.25	1.25	1.25	1.2	1.10	1.2			
20 Margin Sa 9 95 <	19	Active material capacity, mAh/g:		330	330	330	330	330	170	330			
21 Acte material 95	20	Weight %											
bit 0	21	Active material		95	95	95	95	95	89	95			
Ed. Honder 5	22	Carbon		0	0	0	0	0	6	0			
comment water <	23	Binder		5	5	5	5	5	5	5			
Car Vision State Set S4	24	Binder solvent		Water	Water	VVater	Water	Water	VVater	Water			
Constraint 2 24	25	<u>Voia, V01% %</u>		34	34	34	34	34	40	34			
c/r 2/4 2/4 2/4 2/4 2/4 2/4 2/4 2/4 20 Carbon 1.35 1.35 1.35 1.36	26	Density, g/cm [*]		0.07	0.01	0.07	0.04	0.01	0.40	0.04			
20 Jabon 1.39 1.30 1.30 1.30 1.30 1.30 1.30 1.30 1.30 1.30 1.30 1.30	27	Active material		2.24	2.24	2.24	2.24	2.24	3.40	2.24			
22 Dirold 1.10	28	Carbon		1.95	1.95	1.95	1.95	1.95	1.95	1.95			
and Material Aluminum	29	Binder		1.1	1.10	1.10	1.10	1.10	1.10	1.10			
and material Autminum	30	Positive Foll								A1 .			
Jac Deckess, jum ZU ZU <thzu< th=""> ZU <thzu< th=""> <thzu< th=""> <thzu< th=""></thzu<></thzu<></thzu<></thzu<>	31	Material		Aluminum	Aluminum	Aluminum	Aluminum	Aluminum	Aluminum	Aluminum			
31 Agative Fold Copper Coppe	32	Inickness, µm		20	20	20	20	20	20	20			
44 Material Copper Copper <thcopper< th=""> <thcoper< th=""></thcoper<></thcopper<>	33	Negative Foil											
discretes, µm	34	Material		Copper	Copper	Copper	Copper	Copper	Aluminum	Copper			
36 Separator 20	35	l hickness, μm		12	12	12	12	12	20	12			
37 Inickness, µm 20 <td>36</td> <td>Separator</td> <td></td>	36	Separator											
38 Viol. (volte %, % b0 50 50 50 50 50 39 Viol. (volte %, % 0.46	37	Thickness, µm		20	20	20	20	20	20	20			
39 Dentity, g/cm ³ 0.46 0.46<	38	Void, Vol% %		50	50	50	50	50	50	50			
40 Electrolyte density, g/cm ³ 1.2 1.20	39	Density, g/cm [°]		0.46	0.46	0.46	0.46	0.46	0.46	0.46			
11 Cell Voltage and Resistance Parameters n <td>40</td> <td>Electrolyte density, g/cm³</td> <td></td> <td>1.2</td> <td>1.20</td> <td>1.20</td> <td>1.20</td> <td>1.20</td> <td>1.20</td> <td>1.20</td> <td></td> <td></td> <td></td>	40	Electrolyte density, g/cm ³		1.2	1.20	1.20	1.20	1.20	1.20	1.20			
42 OCV at 20% SOC, V 3 826 3 561 3 565 3 576 3 671 3 246 2 408 3 826 3 Open circuit voltage at 50% SOC, V 3 584 3 680 3 750 3 671 3 242 2.514 3 954 44 Solid state diffusion limiting Crate (10-s), A/Ah 120 27 27 120 200 120 45 Negative electrode cm ² /cm ³ 49200 7400 750	41	Cell Voltage and Resistance P	Parameter	5									
43 Open circuit voltage at 50% SOC, V 3 3 3 5 3 671 3.282 2.514 3 3954 44 Solid state diffusion limiting Crate (10-s), A/Ah 120 27 27 27 120 200 120 120 45 Negative electrode cm ² (cm ³) 49200 8900 8900 8900 420000 74000 720 720 72	42	OCV at 20% SOC V		3.826	3.551	3.565	3.516	3.246	2.408	3.826			
44 Solid state diffusion limiting C-rate (10-s), A/Ah 120 27 27 27 120 200 120 45 Negative electrode cm ² /cm ³ 74000 74000 74000 74000 500000 74000 4000 47 Electrode cm ² /cm ³ 49200 8900 8900 8900 42000 49200 49200 48 Selected ASI value 25 -	43	Open circuit voltage at 50% SOC, V		3.954	3.680	3.750	3.671	3.282	2.514	3.954			
45 Negative electrode cm ² /cm ³ 74000 74000 <t< td=""><td>44</td><td>Solid state diffusion limiting C-rate (1</td><td>10-s), A/Ah</td><td>120</td><td>27</td><td>27</td><td>27</td><td>120</td><td>200</td><td>120</td><td></td><td></td><td></td></t<>	44	Solid state diffusion limiting C-rate (1	10-s), A/Ah	120	27	27	27	120	200	120			
46 Positive electrode cm ² /cm ³ 49200 8900 8900 8900 42000 49200 49200 7T Electrode system ASI for power, ohm-cm ² 4 At 50% SOC, 2-sec burst 13 18 21 23.5 20 6 13 45 OK 500% SOC, 2-sec burst 20 23.6 26.6 31 25 8 20 51 At 20% SOC concerction factor 2 3 3 3 1.5 1.5 2 2ASI correction factor 2 3 3 3 1.5 1.5 2	45	Negative electrode cm ² /cm ³		74000	74000	74000	74000	74000	500000	74000			
47 Electrode system ASI for power, ohm-cm ² 25 10	46	Positive electrode cm ² /cm ³		49200	8900	8900	8900	420000	49200	49200			
48 Selected ASI value 25 10 10 10 10 49 At 50% SOC, 2-see burst 13 18 21 23.5 20 6 13 51 At 20% SOC, 10-see burst 25 30 33 36 32 9.4 25 2 ASI correction factor 2 3 3 3 1.5 1.5 2 2 ASI correction factor 2 3 3 3 1.5 1.5 2 54 Electrode system ASI for energy, ohm-cm ² 44 51.9 58.5 68.2 55.0 11.7.6 44.0 55 Maximum allowable ASI for brenergy, of total 70 100	47	Electrode system ASI for power, ohr	m-cm ²										
49 At 50% SOC, 2-sec burst 13 18 21 23.5 20 6 13 50 At 50% SOC, 10-sec burst 20 23.6 26.6 31 25 8 20 51 At 20% SOC, 10-sec burst 25 30 33 36 32 9.4 25 25 53 Maximum allovable ASI for limiting capacity 37.5 -	48	Selected ASI value		25									
50 At 50% SOC, 10-sec burst 20 23 6 26 6 31 25 8 20 51 At 20% SOC range, 10-sec burst 25 30 33 36 32 9.4 25 53 Maximum allowable ASI for limiting capacity 37.5 3 1.5 1.5 2 1.6 54 Electrode system ASI for energy, 0hm-cm ² 44 51.9 58.5 68.2 55.0 11.76 44.0 51.9 56 Maximum electrode coating thickness, µm 100 11.5	49	At 50% SOC, 2-sec burst		13	18	21	23.5	20	6	13			
51 At 20% SOC range, 10-sec burst 25 30 33 36 32 9.4 25 52 ASI correction factor 2 3 3 3 1.5 1.5 2 54 Electrode system ASI for inmiting capacity 37.5 - - - - - - 54 Electrode coating thickness, µm 100	50	At 50% SOC, 10-sec burst		20	23.6	26.6	31	25	8	20			
	51	At 20% SOC range, 10-sec burst	t	25	30	33	36	32	9.4	25			
53 Maximum allowable ASI for limiting capacity 37.5 Image: Constraint of Constraints of Constrain	52	ASI correction factor		2	3	3	3	1.5	1.5	2			
54 Electrode system ASI for energy, ohm-cm ² 44 51.9 58.5 68.2 55.0 11.76 44.0 56 Maximum electrode coating thickness, µm 1000 1000 1000 1000 1000 1000 1000 1000 1000 1000 1000 1000 1000 1000 1000 1000 1000 1000	53	Maximum allowable ASI for limiting of	capacity	37.5									
55 Maximum electrode coating thickness, μm 100	54	Electrode system ASI for energy, oh	m-cm ²	44	51.9	58.5	68.2	55.0	11.76	44.0			
56 Available battery energy, % of total 70 6 7 8 8 7 70 <t< td=""><td>55</td><td>Maximum electrode coating thicknes</td><td>ss, μm</td><td>100</td><td>100</td><td>100</td><td>100</td><td>100</td><td>100</td><td>100</td><td></td><td></td><td></td></t<>	55	Maximum electrode coating thicknes	ss, μm	100	100	100	100	100	100	100			
57 Selected % energy microHEV and HEV-HP 70	56	Available battery energy, % of	total										
56 microHEV and HEV-HP 25 <td>57</td> <td>Selected % energy</td> <td></td> <td>70</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>	57	Selected % energy		70									
59 PHEV 70 70 70 70 70 75 70 60 EV 85 85 85 85 90 85 90 85 85 90 85 85 90 85 60 Cell Materials Costs Image: Costs Imag	58	microHEV and HEV-HP		25	25	25	25	25	25	25			
60 EV 85 85 85 85 85 90 85 61 Cell Materials Costs r NCA-G NMC441-G NMC333-G LFP-G LMO-LTO LMO-G Override Values 62 Positive Electrode, \$/kg p Cost p Cost p 64 Carbon Black 6.80 100 6.80 <t< td=""><td>59</td><td>PHEV</td><td></td><td>70</td><td>70</td><td>70</td><td>70</td><td>70</td><td>75</td><td>70</td><td></td><td></td><td></td></t<>	59	PHEV		70	70	70	70	70	75	70			
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	60	EV		85	85	85	85	85	90	85			
62 Positive Electrode, \$/kg p Cost p 63 Active material 10:00 0:95 33:00 26:00 31:00 20:00 10:00 10:00 64 Carbon Black 6:80 100 6:80	61	Cell Materials Costs			NCA-G	NMC441-G	NMC333-G	LFP-G	LMO-LTO	LMO-G	Overrid	e Values	
63 Active material 10:00 0:95 33:00 26:00 31:00 20:00 10:00 10:00 64 Carbon Black 6:80 100 6:80	62	Positive Electrode, \$/kg		р							Cost	р	
64 Carbon Black § 80 100 6.80 6.80 6.80 6.80 6.80 65 Binder PVDF 10.00 10.0	63	Active material	10.00	0.95	33.00	26.00	31.00	20.00	10.00	10.00			
65 Binder PVDF 10.00	64	Carbon Black	6.80	1.00	6.80	6.80	6.80	6.80	6.80	6.80			
66 Binder Solvent (NMP) 3.20 <td>65</td> <td>Binder PVDF</td> <td>10.00</td> <td>1.00</td> <td>10.00</td> <td>10.00</td> <td>10.00</td> <td>10.00</td> <td>10.00</td> <td>10.00</td> <td></td> <td></td> <td></td>	65	Binder PVDF	10.00	1.00	10.00	10.00	10.00	10.00	10.00	10.00			
67 Negative electroide material, \$/kg 19.00 10.00 10.00 10.00 10.00 10.00 10.00 10.00 10.00 10.00 10.00 10.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 <td>66</td> <td>Binder Solvent (NMP)</td> <td>3.20</td> <td>1.00</td> <td>3.20</td> <td>3.20</td> <td>3.20</td> <td>3.20</td> <td>3.20</td> <td>3.20</td> <td></td> <td></td> <td></td>	66	Binder Solvent (NMP)	3.20	1.00	3.20	3.20	3.20	3.20	3.20	3.20			
66 Active Material 19:00 10:00 10:00 10:00 10:00 10:00 10:00 10:00 10:00 10:00 10:00 10:00 10:00 0:00	67	Negative electrode material, \$/kg											
69 Carbon Black 5.80 1.00 6.80 6.80 6.80 6.80 6.80 6.80 70 Binder 10.00 <td>68</td> <td>Active Material</td> <td>19.00</td> <td>1.00</td> <td>19.00</td> <td>19.00</td> <td>19.00</td> <td>19.00</td> <td>12.00</td> <td>19.00</td> <td></td> <td></td> <td></td>	68	Active Material	19.00	1.00	19.00	19.00	19.00	19.00	12.00	19.00			
Dinder Dinder <thdin< th=""> Dinder Dinder</thdin<>	69	Carbon Black	6.80	1.00	6.80	6.80	6.80	6.80	6.80	6.80			
Dinder Solvent 0.00 0.00 0.00 0.00 0.00 0.00 0.00 72 Positive current collector foil, \$/m² 0.80 1.00 0.80 1.80 7.80 7.80 7.00 2.00<	70	Binder Divert	10.00	1.00	10.00	10.00	10.00	10.00	10.00	10.00			
72 Positive current collector foil, \$/m ² 0.80 1.80 0.80 0.80 0.80 0.80 0.80 73 Negative current collector foil, \$/m ² 1.80 1.80 1.80 1.80 1.80 1.80 74 Separators, \$/m ² 2.00 1.00 2.00 2.00 2.00 2.00 2.00 75 Electrolyte, \$/L 21.60 1.00 21.60 21.60 21.60 21.60 21.60 21.60	/1	Binder Solvent	0.00	1.00	0.00	0.00	0.00	0.00	0.00	0.00			
73 Negative current collector foil, \$/m² 1.80 1.80 1.80 1.80 0.80 1.80 74 Separators, \$/m² 2.00 1.00 2.00 2.00 2.00 2.00 2.00 2.00 7.00 2.00 2.00 2.00 2.00 7.00 2.00 7.00 2.00 7.00 2.00 7.00<	12	Positive current collector foil, \$/m ²	0:80	1.00	0.80	0.80	0.80	0.80	0.80	0.80			
74 Separators, \$/m ⁴ 2.00 2.00	73	Negative current collector foil, \$/m ²	1.80	1.00	1.80	1.80	1.80	1.80	0.80	1.80			
75 Electrolyte, 5/L 21.60 21.60 21.60 21.60 21.60 76	74	Separators, \$/m ²	2.00	1.00	2.00	2.00	2.00	2.00	2.00	2.00			
/6	75	Electrolyte, \$/L	21.60	1.00	21.60	21.60	21.60	21.60	21.60	21.60			
	76												

Figure 6.3 Chem worksheet

	3 - ((≃ - =				BatPaC v	2.1 - Microso	ft Excel						×
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2					wanga	Rottony 1	Bottony 2	Rettory 2	Potton: 4	Potton 5	Patton 6	Pottony 7		
3	Parame	ters for Fin	ished Cell			Dattery 1	Dattery 2	Dattery 5	Dattery 4	Dattery 5	Dattery 0	Dattery r		
4	Positivo	Electrode		Woight %	Density									
6	Active mat	torial	9	89	4 23	106.03	159 44	212.60	265 77	318 95	372 14	425 33		
7	Carbon	teriar		6	1.825	7.15	10.75	14.33	17.92	21.50	25.09	28.67		
8	Binder			5	1.77	5.96	8.96	11.94	14.93	17.92	20.91	23.89		
9	Void	Vol. %	32			-	-	-	-	-	-	-		
10	Total			100	2.504	119.14	179.15	238.88	298.62	358.37	418.13	477.90		
11	Negative	e Electrode,	g	Weight %	Density	10.10	50.04	70.57				157.00		
12	Active mat	terial		95	2.24	40.10	59.94	79.57	99.17	118.75	138.31	157.86		
14	Binder			5	1.55	2 11	3 15	4 19	5.22	6.25	7.28	8.31		
15	Void	Vol. %	34			-	-	-	-	-	-	-		
16	Total			100	1.406	42.22	63.09	83.76	104.39	125.00	145.59	166.17		
17	Balance	of Cell		Thick., µm	Density									
18	Positive fo	il, m²	AI	20	2.70	0.357	0.382	0.505	0.627	0.750	0.872	0.993		
19	Negative f	oil, m ²	Cu	12	8.92	0.387	0.415	0.547	0.677	0.808	0.938	1.067		
20	Separator	, m ²		20	0.46	0.723	0.770	1.017	1.263	1.508	1.753	1.998		
21	Electrolyte	e, L			1.20	0.0348	0.0489	0.0650	0.0811	0.0972	0.1132	0.1293		
22	Positive te	erminal assemb	bly, g			4.1	5.0	5.9	6.6 21.0	7.3	7.9	8.5		
23	Thickness	of cell contain	ory, g er aluminum lave	er um		100	10.5	19.4	21.9	24.2	20.2	20.2		
25	Thickness	of cell contain	er (PET-AI-PP).	μm		150	150	150	150	150	150	150		
26	Density of	cell container,	g/cm3			2.2	2.2	2.2	2.2	2.2	2.2	2.2		
27	Cell conta	iner (PET-AI-P	P), g			13.5	17.9	22.8	27.6	32.4	37.1	41.8		
28	Cell mass	, g				302	412	544	674	805	934	1,064		
29	Length-to-	width ratio for p	positive electrode	9		3.00	3.00	3.00	3.00	3.00	3.00	3.00		
31	Thickness	of cell edge fr	om positive elect	rode to outside	of fold mm	0.0	0.0	0.0	0.0	0.0	0.0	0.0		
32	Thickness	of terminal ma	aterial, mm		or loid, min	1.00	1.00	1.00	1.00	1.00	1.00	1.00		
33	Top of pos	sitive electrode	to top of termina	l, mm		15	15	15	15	15	15	15		
34	Cell Cap	acity Param	eters											
35	Positive a	ctive material c	apacity, mAh/g:			100	100	100	100	100	100	100		
36	Positive el	lectrode capaci	ity, Ah/cm ³			0.223	0.223	0.223	0.223	0.223	0.223	0.223		
37	Negative a	active material of	capacity, mAh/g:			330	330	330	330	330	330	330		
38	Negative e	electrode capac	ity, Ah/cm ³			0.441	0.441	0.441	0.441	0.441	0.441	0.441		
39	Negative-t	o-positive capa	icity ratio after for	rmation		1.20	1.20	1.20	1.20	1.20	1.20	1.20		
40	Cell Volt	age and Res	sistance Parar	meters		2 000	2 026	2 000	2 0.00	2 000	2 000	2 000		
41	Open circi	uit voltage aver	age for discharge	v		3.954	3 954	3 954	3.954	3.954	3 954	3.954		
43	Electrode	system ASI for	r enerav, ohm-cn	n ²		44.0	44.0	44.0	44.0	44.0	44.0	44.0		
44	Excess ne	egative area, %	3,			4.01	3.38	2.93	2.62	2.39	2.21	2.07		
45	Default ma	aximum allowa	ble electrode coa	ating thickness	, μm	100	100	100	100	100	100	100		
46	Cell termir	nal contact volt	age loss, % of c	ell OCV		0.01	0.01	0.01	0.01	0.01	0.01	0.01		
47	Rate of ter	rminal tempera	ture rise at full po	ower, °C/sec		0.05	0.05	0.05	0.05	0.05	0.05	0.05		
48	arget %	UCV at full pov	ver isted for thickness	e limit		80.0	81 S	87.0	80 8	8U 91 5	8U 92.7	93.6		<u> </u>
50	Batten	Input Parar	neters	55 mm		00.0	01.0	01.0	03.0	51.5	32.1	55.0		
50	Vehicle	type (micro		PHEV EV		DHEV		DHEV	DHEV	DHEV	DHEV	DHEV		
52	Pack heat	transfer fluid /	EG-W CA Cool	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		EG-W	FG-W	FG-W	FILEV FG-W	FG-W	EG-W	EG-W		
53	Duration o	f power burst (10 or 2), s	,		10	10	10	10	10	10	10		
54	Battery pa	ck power, kW				60	60	60	60	60	60	60		
55	Number of	f cells per mod	ule			24	24	24	24	24	24	24		
56	Number of	f cells in paralle	el			1	1	1	1	1	1	1		
5/	Number of	r modules in ro	W			4	4	4	4	4	4	4		
59	Number of	f modules ner h	es per pack			4	4	4	4	4	4	4		
60	Number of	f modules in pa	rallel			1	1	1	1	1	1	1		
61	Number of	f packs per veh	icle (parallel or s	eries)		1	1	1	1	1	1	1		
62	Paralle	el packs (P) or	series (S)											
63	Cells per l	battery pack				96	96	96	96	96	96	96		
64	Voltage dr	per pattery sy	stem for packe with or	a row of modu	les V	96	96	96	96	96	96	96		
66	Battery pa	op ior bus par ack insulation t	hickness. mm		ica, v	10	10	10	10	10	10	10		
67	Battery ja	cket total thick	ness, mm			12	12	13	13	13	14	14		
68	Number o	of batteries m	anufactured pe	r year		100,000	100,000	100,000	100,000	100,000	100,000	100,000		
69	Cell Cher	nistry Input												
70	Battery P	erformance a	nd Design Inpu	t										
71	Calcula	ted Cell Par	rameters											

Figure 6.4 Top portion of Battery Design worksheet.

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50	Battery I	Input	Parame	eters											_
51	Vehicle t	type (m	hicroHE	V. HEV-HE	P. PHEV. EV)	PHEV	PHEV	PHEV	PHEV	PHEV	PHEV	PHEV		
52	Pack heat	transfer	r fluid (EG	-W, CA, Co	olA)	,	EG-W	EG-W	EG-W	EG-W	EG-W	EG-W	EG-W		
53	Duration of	f power	burst (10	or 2), s	,		10	10	10	10	10	10	10		
54	Battery par	ick powe	er, kW				60	60	60	60	60	60	60		
55	Number of	cells pe	er module	•			24	24	24	24	24	24	24		
56	Number of	cells in	parallel				1	1	1	1	1	1	1		
57	Number of	module	s in row				4	4	4	4	4	4	4		
58	Number of	rows of	modules	per pack			1	1	1	1	1	1	1		
59	Number of	module	s per bat	tery pack			4	4	4	4	4	4	4		
60	Number of	module	s in para	llel			1	1	1	1	1	1	1		
61	Number of	packs	per vehicl	e (parallel or	series)		1	1	1	1	1	1	1		
62	Parallel	l packs	(P) or se	ries (S)											
63	Cells per b	pattery p	ack				96	96	96	96	96	96	96		
64	Total cells	per bat	tery syste	em			96	96	96	96	96	96	96		
05	Voltage dro	op for bl	us bar tor	packs with o	one row of mod	lies, v	0.03	0.03	0.03	0.03	0.03	0.03	0.03		
67	Dattery par	ick insul	ation thickne	ckness, mm			10	10	10	10	10	14	10		
07	Dattery jac		al thickne	ss, mm			12	12	13	13	13	14	14		
60	Cell Chem	n Datter	nes man	ulactureu p	ei yeai		100,000	100,000	100,000	100,000	100,000	100,000	100,000		
70	Pottony Dr	nisuy ii	iput maa and	Design Inn											
70	Calculat			n Design inp	au contraction of the second se										
/1	calculat	iea Ce	iii marai	meters											
72	Capacity,	, Ah													
73	Cell group	capacit	у				10.6	15.9	21.3	26.6	31.9	37.2	42.5		
74	Cell capac	ity					10.6	15.9	21.3	26.6	31.9	37.2	42.5		
75	ASI Calcu	ulation	1												
76	Limiting cu	urrent de	ensity, m	A/cm ²			85	85	85	85	85	85	85		
77	Limiting C-	-rate, A/	Ah				120	120	120	120	120	120	120		
78	Electrode s	system	ASI for p	ower at SOC	for vehicle type	e, ohm-cm ²	23.77	23.54	23.51	23.49	23.49	23.48	23.48		
79	Current col	llector r	esistance	e parameter,	ohms		0.005409	0.005409	0.005409	0.005409	0.005409	0.005409	0.005409		
80	Current col	llector A	ASI, ohms	s-cm ²			0.901	1.227	1.599	1.967	2.333	2.697	3.059		
81	Cell termin	nal and o	connectio	n ASI, ohms	-cm ²		0.126	0.112	0.128	0.141	0.154	0.165	0.176		
82	Resistance	e of mod	dule and	pack per moo	dule, ohms		0.0000616	0.0000630	0.0000670	0.0000691	0.0000705	0.0000714	0.0000721		
83	Resistance	e of mod	dule and p	pack hardwar	re per cell, ohm	s	0.0000026	0.0000026	0.0000028	0.0000029	0.0000029	0.000030	0.0000030		
84	Total cell h	hardware	e and bat	tery ASI, ohn	n-cm ²		1.044	1.358	1.753	2.143	2.529	2.912	3.292		
85	Total cell A	ASI for p	ower, oh	m-cm ²			24.81	24.90	25.26	25.64	26.02	26.39	26.77		
86	Total cell A	ASI for e	energy (C	/3 rate), ohm	-cm ²		45.68	45.81	46.21	46.60	46.99	47.37	47.75		
87	Electrod	e Coat	tina Thi	ckness Ca	lculation										
88	Negative-to	o-positiv	e electro	de thickness	ratio		0.607	0.607	0.607	0.607	0.607	0.607	0.607		
89	Positive ele	ectrode	thicknes	s, if at maxin	num . um		100.0	100.0	100.0	100.0	100.0	100.0	100.0		
90	Negative el	lectrode	thicknes	s, if at maxi	mum , um		60.7	60.7	60.7	60.7	60.7	60.7	60.7		
91	Thickest el	lectrode	e, μm				100.0	100.0	100.0	100.0	100.0	100.0	100.0		
92	Positive ele	ectrode	thicknes	s parameter,	μm		71.8	107.7	141.5	174.3	206.1	237.1	267.1		
93	Negative el	lectrode	thicknes	s parameter	, μm		43.6	65.4	85.9	105.8	125.1	143.9	162.1		
94	Positive ele	ectrode	thicknes	s at adjusted	I % OCV, μm		71.8	100.0	100.0	100.0	100.0	100.0	100.0		
95	Negative el	lectrode	thicknes	ss at adjusted	d % OCV, μm		43.6	60.7	60.7	60.7	60.7	60.7	60.7		
96	Cell Area	a Calcu	lation.	cm ²											
97	Area deter	mined a	t target 9	% OCV			6.621	6.643	6,741	6.841	6.942	7.043	7,144		
98	Area limite	ed by ma	ax. allowe	ed electrode t	thickness,		4,757	7,153	9,539	11,924	14,310	16,696	19,083		
99	Cell area b	pased or	n total AS	SI for power, o	cm ²		6.621	7,153	9.539	11,924	14.310	16.696	19.083		=
100	Cell Dime	ension	IS I	Person (- 1041	.,	-,	,•= (,			
101	Number of	bicell	vers (97	% nacking de	ensity)		25	19	19	19	19	19	19		
102	Width of pr	ositive e	lectrode	mm			67	79	92	102	112	121	130		
103	Length of r	positive	electrode	, mm			201	238	275	307	337	363	389		
104	Length of c	current o	collector t	tabs, mm			16	16	16	16	16	16	16		
105	Width of te	erminals	, mm				59	71	84	94	104	113	122		
106	Length of t	terminal	material,	mm			26	26	26	26	26	26	26		
107	Width of ce	ell, mm					69	81	94	104	114	123	132		
108	Length of c	cell, mm	1				231	268	305	337	367	393	419		
109	Volume of	cell, cm	n ³				127	174	228	282	335	388	440		
110	Module	Paran	neters												
111	Module car	pacity.	Ah				10.6	15.9	21.3	26.6	31.9	37.2	42.5		
112	Mass of ea	ach cell	group int	erconnect (c	opper), g		0.0	0.0	0.0	0.0	0.0	0.0	0.0		
113	Module sta	ate-of-ch	narge regi	ulator assem	bly, g		192	192	192	192	192	192	192		
114	Terminal h	eating fa	actor, W/	g	_		0.019	0.019	0.019	0.019	0.019	0.019	0.019		
115	Terminal re	esistanc	e factor,	A-ohms/cm			0.00054	0.00054	0.00054	0.00054	0.00054	0.00054	0.00054		
116	Module ter	rminals,	if more t	han one mod	ule (each 2.0-ci	m long), g	27	27	25	24	24	23	23		
117	Module ter	rminal re	esistance	both termina	als, ohms		0.0000105	0.0000107	0.0000114	0.0000118	0.0000120	0.0000122	0.0000123		
118	Module res	sistance	, ohms				0.0014890	0.0015233	0.0016194	0.0016709	0.0017034	0.0017258	0.0017422		
119	Module wa	all mater	ial				aluminum	aluminum	aluminum	aluminum	aluminum	aluminum	aluminum		
120	Module wa	all thickr	ness, mm	1			0.5	0.5	0.5	0.5	0.5	0.5	0.5		
121	Length of a	aluminu	m conduc	ctor or therma	al enclosures, n	nm	201	238	275	307	337	363	389		
122	Tetel	ot alum	inum con	auctor or the	ermai enclosure	, mm	0.40	0.40	0.40	0.40	0.40	0.40	0.40		
123	i otal mass	s of alun	ninum co	nductors or t	nermai enclosu	res, g	461	625	813	998	1,183	1,366	1,548		(
124	Datance of Module les	i module	e material	is, g			21/	262	308	351	391	430	46/		
125	Module with	igin, mr	n				233	2/0	307	211	309	395	421		
120	Modulo koi	iaht m	n				211	211	211	211	116	105	122		
127	Module riel	agric, mr lume 1					3.48	4 73	6 17	7.60	9.02	10.43	11.83		
120	Module ma	ass ko					8 14	11.00	14 39	17.75	21 10	24 44	27 77		
.20							9.14				21.19		2000		

Figure 6.5 Middle portion of Battery Design worksheet

The Battery Design worksheet automatically receives input from the Chem worksheet. These values are shown in purple (Figures 6.4 and 6.6) and must not be altered on the Battery Design worksheet. As explained above, cell chemistry values may be adjusted on the Chem worksheet. The operator provides battery design input in the aqua colored cells (Figures 6.4 and 6.6). The battery input parameters on lines 54 to 62 (Figure 6.4) and lines 179 to 181 (Figure 6.6) are the only input values that the operator is required to provide to study a group of batteries. The type of vehicle battery (microHEV, HEV-HP, PHEV, or EV) on line 51 in Figure 6.4, is another important variable to be specified. One performs the selection by typing the name of the vehicle battery type in cell F51. While the correct spelling is important, capitalization is not. This selection automatically determines the state of charge at which full power is designated (thus, the open-circuit voltage and ASI for full power) and the length of the power burst (2 seconds for microHEV and 10 seconds for all others). It is expected that the majority of the remaining default values should serve well for most batteries; however, the user may also change to their exact specifications. The thermal management approach is selected on line 52 in Figure 6.4. One can select between a thermal management approach using liquid (EG-W), cabin air (CA), or conditioned cabin air (CoolA). The different module and battery formats for air and liquid cooling are displayed in later worksheets of BatPaC.

The cell capacity (lines 179 to 181 in Figure 6.6) can be set in any of three ways: (1) directly specifying the capacity (Ah) on line 179, (2) specifying the total battery energy on line 180 or (3) specifying the electric range of the vehicle (miles). Only one of the three lines should be filled in and the others should be blank. The model will follow the directions of the top-most line with non-zero values.

The number of batteries manufactured per year is selected on line 68 in Figure 6.4. Changing this value from the default value of 100,000, which is the manufacturing rate for the baseline plant, will change the manufacturing cost.

If it is desired to study more than five batteries in the same workbook it is only necessary to add additional columns by copying the battery 7 column to the right as many times as desired. Care should be taken that the appropriate values are maintained when the cells are copied over. The aqua colored cells are typically the source of any problems. The same column additions must also be done for all other worksheets containing calculations.

6.2.4 Remaining Worksheets

The cost calculations are done on the Manufacturing Cost worksheet and the results for the model are shown on the Summary of Results worksheet (Figure 6.7). No parameters need to be entered on these worksheets by the operator; all of the input for these worksheets is from the Battery Design and the Cost Input worksheets. Tables for presentations or for preparing graphs of the data can be assembled at the bottom of either the Battery Design or the Summary of Results worksheet. These tables can be transferred to a blank worksheet for more complex studies. For instance, results for different cell chemistries can be copied and pasted (special paste, values and numbers formats) to a blank worksheet. On the last worksheets, the cell, module, and battery design, as well as the baseline plant are sketched.

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	A	В	С	D	E	F	G	H		J	K	L	М	
110	Module Pa	arameters												
111	Module capa	city, Ah				10.6	15.9	21.3	26.6	31.9	37.2	42.5		
112	Mass of each	n cell group ir	nterconnect (co	opper), g		0.0	0.0	0.0	0.0	0.0	0.0	0.0		
113	Module state	-of-charge re	gulator assemi	bly, g		192	192	192	192	192	192	192		
114	Terminal hea	ting factor, W	//g			0.019	0.019	0.019	0.019	0.019	0.019	0.019		
115	Terminal resi	stance factor	, A-ohms/cm	1. (0.00054	0.00054	0.00054	0.00054	0.00054	0.00054	0.00054		
116	Module term	nais, ir more	than one mod	ule (each 2.0-c	m long), g	21	21	25	24	24	23	23		
117	Module termi	nai resistanc	e both termina	is, onms		0.0000105	0.0000107	0.0000114	0.0000118	0.0000120	0.0000122	0.0000123		
110	Module resis	natorial				0.0014690	0.0015233	0.0010194	0.0010709	0.0017034	0.0017250	0.0017422		
120	Module wall t	hicknose m	m			0.5	0.5	0.5	0.5	0.5	0.5	0.5		
121	Length of alu	minum condu	ictor or therms	l enclosures r	nm	201	238	275	307	337	363	389		
122	Thickness of	aluminum co	inductor or the	rmal enclosure	mm	0.40	0.40	0.40	0.40	0.40	0.40	0.40		
123	Total mass o	f aluminum c	onductors or th	hermal enclosu	ires a	461	625	813	998	1 183	1.366	1.548		
124	Balance of m	odule materia	als. o			217	262	308	351	391	430	467		
125	Module lengt	h, mm				233	270	307	339	369	395	421		
126	Module width	, mm				211	211	211	211	211	211	211		
127	Module heigh	nt, mm				71	83	95	106	116	125	133		
128	Module volun	ne, L				3.48	4.73	6.17	7.60	9.02	10.43	11.83		
129	Module mass	s, kg				8.14	11.00	14.39	17.75	21.10	24.44	27.77		
130	Calculate	d Battery I	Pack Paran	neters										
131	Battery pack	capacity, Ah	1			10.6	15.9	21.3	26.6	31.9	37.2	42.5		
132	Total battery	pack energy	storage, kWh			4.00	6.00	8.00	10.00	12.00	14.00	16.00		
133	Useable batt	ery energy st	orage, kWh			2.80	4.20	5.60	7.00	8.40	9.80	11.20		
134	OCV at full p	ower, V				367.3	367.3	367.3	367.3	367.3	367.3	367.3		
135	Nominal batt	ery voltage (C	OCV at 50% S	DC)		379.6	379.6	379.6	379.6	379.6	379.6	379.6		
136	Battery powe	r at target %	UCV and SO	J, KW		60.0	64.6	84.9	104.6	123.7	142.2	160.3		
137	Maximum cu	rrent at full p	ower, A			204	200	188	182	178	176	1/5		
138	Maximum cu	rrent density	at full power, r	mA/cm⁴		30.84	27.90	19.68	15.26	12.47	10.55	9.15		
139	C-rate at full	power, A/Ah	halanna 111			19.3	12.5	8.8	6.8	5.6	4.7	4.1		
140	Coolant space	e above and	perow modules	s, mm		6.0	5.5	4.8	4.3	3.9	3.6	3.4		
141	Inickness of	module com	pression plate:	s (steel), mm		1.5	1.5	1.5	1.5	1.5	1.5	1.5		
142	Battery pack	length (A dir	mension), mm			000	8//	8/8	8//	8//	8/9	8/8		
143	Batteny pack	wath (B ann	nension), mm			205	302	341	373	403	431	457		
144	Length of pack	k extensions	for air cooled	nacks mm		50	50	50	50	50	50	100 50		
146	Thickness of	nack extensions	ions for air-cor	led nacks mn	1	0	0	0	0	0	0	0		
147	Total volume	of pack exter	nsions for air-c	ooled packs. L		0.0	0.0	0.0	0.0	0.0	0.0	0.0		
148	Pack integrat	tion unit (BM	S & disconnec	ts), L		4.0	4.0	4.0	4.0	4.0	4.0	4.0		
149	Volume of ba	ttery pack ar	nd integration u	init, L		28.8	35.3	43.2	50.1	56.9	64.7	71.4		
150	Mass of each	n module inte	r-connect (5-ci	m long), g		34	33	31	30	30	29	29		
151	Mass of mod	ule compress	sion plates and	l steel straps,	g	466	612	773	932	1091	1248	1406		
152	Mass of bus	bar for packs	with one row	of modules, g		736	719	675	653	640	632	625		
153	Resistance of	f module inte	rconnects if m	ore than one n	nodule, ohms	0.0000472	0.0000483	0.0000513	0.0000530	0.0000540	0.0000547	0.0000552		
154	Resistance of	f battery pac	k terminals			0.0000157	0.0000161	0.00001/1	0.00001//	0.0000180	0.0000182	0.0000184		
155	Power of batt	ery neaters,	KVV	V) ka		2.0	2.0	2.0	2.0	2.0	2.0	2.0		
150	Battony cools	ery pack near	un inckot ka	v), kg		2.70	2.91	2.87	2.86	2.84	2.84	2.85		
157	Batteny coola	t mass with	motor g/om ²			0.57	0.57	0.94	2.05	0.94	1.11	2.05		
150	Total mass of	f nack ovton	nieter, g/cm	olod packe, ka		0.07	0.07	0.04	0.04	0.04	0.00	0.00		
160	Battery jacke	t mass ko	sions for all co	ored packs, kg		5.1	5.8	8.7	9.6	10.4	14.2	15.2		
161	Pack integral	tion unit (BM	S & disconnec	ts. ave. densit	v = 1.0), kg	4.0	4.0	4.0	4.0	4.0	4.0	4.0		
162	Mass of batte	ery pack and	integration uni	t, ka		44.4	56.7	73.2	87.5	101.7	118.8	133.1		
163	Battery syste	em volume (a	II packs), L			28.8	35.3	43.2	50.1	56.9	64.7	71.4		
164	Battery syste	em mass (all	packs), kg			44.4	56.7	73.2	87.5	101.7	118.8	133.1		
165	Battery Co	ooling Sys	stem											
166	Heat generat	ion rate for ba	attery system.	w		1326	1215	883	697	577	493	432		
167	Cooling Syst	em				Refrig 3	Refrig 3	Refrig 3	Refrig 3	Refrig 2	Refrig 2	Refrig 2		
168	Mass and Vo	olume of Cool	ing System Ex	terior to Batte	ry Packs									
169	Mass, kg					5.0	5.0	5.0	5.0	3.0	3.0	3.0		
170	Volume, I					2.0	2.0	2.0	2.0	1.2	1.2	1.2		
171	Vehicle E	ectric Rar	nge											
172	Energy requi	rement of veh	icle, Wh/mile			250.0	250.0	250.0	250.0	250.0	250.0	250.0		
173	Energy requi	rement of bat	tery pack, Wh	/mile		250.0	250.0	250.0	250.0	250.0	250.0	250.0		
174	Constant spe	ed sustained	l at energy req	uirement, mph		58	58	58	58	58	58	58		
175	Available bat	tery energy,	% of total			70	70	70	70	70	70	70		+
176	Vehicle elect	ric range, mil	les			11.20	16.80	22.40	28.00	33.60	39.20	44.80		+
177	Pack Cap	acity Calc	ulation											
178	Select capa	city, battery	energy, or v	ehicle range,	but only one.									
179	Pack capa	acity (Ah)				10			40.0	42.0		40.0		
100	Pack ener	gy (kvVh)				4.0	6.0	8.0	10.0	12.0	14.0	16.0		+
101	Canacity act	imating para	notor			10.5	16.9	21.1	26.3	31.6	36.0	12.2		+
183	Capacity est	C/3 Ah	notor			10.603	15 944	21 260	26 577	31.895	37 214	42 533		+
184	Capacity at C	dina				10.603	15 944	21.200	26.577	31 895	37 214	42.533		+
185	Positive elect	trode thickne	ss			71.8	100.0	100.0	100.0	100.0	100.0	100.0		
186	Positive elect	trode thickne	ss holding			71.8	100.0	100.0	100.0	100.0	100.0	100.0		
187	Convergence	parameter	3			0.30	0.30	0.30	0.30	0.30	0.30	0.30		
188				F	Restart (0/1)	1	1	1	1	1	1	1		
189														
190				Batterv Svs	tem Values									
191				, .,.	Power, kW	60.0	60.0	60.0	60.0	60.0	60.0	60.0		
192					Energy, kWh	4.00	6.00	8.00	10.00	12.00	14.00	16.00		
193					Mass, ko	44.4	56.7	73.2	87.5	101.7	118.8	133.1		
194					Volume, L	28.8	35.3	43.2	50.1	56.9	64.7	71.4		
195					Wh/kg	90	106	109	114	118	118	120		
196					Wh/Ľ	139	170	185	200	211	216	224		
197					\$/kWh	576	423	360	320	293	275	259		
198			Tot	al Battery Cos	t to OEM, US\$	2306	2536	2879	3199	3513	3844	4151		
199			Maximum Tot	al Battery Cos	t to OEM, US\$	2715	3218	3714	4182	4640	5116	5564		+
200			Minimum Tot	al Battery Cos	t to OEM, US\$	2075	2252	2427	2653	2896	3157	3395		+

Figure 6.6 Bottom portion of Battery Design worksheet

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F	File Home Insert	Page	e Layout	Formulas	Data Revi	ew View	Add-Ins						∞ 🕜	- @ X
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	AD	U	D	E	Summa		ulte		J	ĸ	L	IVI	IN	
1					Mongonood	Spinel/C	uits ranhita							
2					Batton 1	Batton 2	Battony 3	Batton A	Batton 5	Batton 6	Battony 7			
4	Calculated Battern	v Para	metere		Dattery	Dattery 2	Dattery J	Dattery 4	Dattery J	Dattery	Dattery			
5	Vehicle electric range	y i ara miles	metera		11.2	16.8	22.4	28.0	33.6	39.2	44.8			
6	Number of battery pack	s			1	10.0	1	20.0	1	1	1			
7	Packs in series or p	parallel												
8	Number of cells per pac	:k			96	96	96	96	96	96	96			
9	Battery system total en	nergy sto	orage, kWh		4.0	6.0	8.0	10.0	12.0	14.0	16.0			
10	Cell capacity, Ah				10.6	15.9	21.3	26.6	31.9	37.2	42.5			
11	Cell group capacity, An				10.6	15.9	21.3	26.6	31.9	31.2	42.5			
12	Pack capacity, An				10.6	15.9	21.3	20.0	31.9	37.2	42.5			
14	Battery system capacit	v. Ah			10.6	15.9	21.3	26.6	31.9	37.2	42.5			
15	Nominal battery system	n voltage	OCV at 50	% SOC),V	380	380	380	380	380	380	380			
16	Battery system power a	at target	% OCV, kV	V	60.0	64.6	84.9	104.6	123.7	142.2	160.3			
17	Required battery syster	m power	, kW		60.0	60.0	60.0	60.0	60.0	60.0	60.0			
18	Target % OCV at full po	ower			80.0	80.0	80.0	80.0	80.0	80.0	80.0			
19	% OCV at full power ad	ljusted fo	or thickness	limit	0.08	81.8	87.0	89.8	91.5	92.7	93.6			
20	Battery system mace //	all packs	no), L a) ka		20.0 ////	35.3 56.7	43.2	50.1 97.4	56.9	04./ 118.9	133.1			
22	Cooling system power r	requirem	ent, W		531	487	354	279	231	198	173			
23	Investment Costs						004	2,5	201	100				
24	Capital equipment cost	includin	g installatio	n, mil\$	133	143	154	165	175	184	192			
25	Building, Land and Utilit	ties	0											
26	Area, m ²				15,917	16,976	18,418	19,721	20,924	22,050	23,115			
27	Cost, \$/m ²				3,000	3,000	3,000	3,000	3,000	3,000	3,000			
28	Building investment,	, mil\$			47.8	50.9	55.3	59.2	62.8	66.1	69.3			
29	Launch Costs													
30	Rate: 5% of direct a	innual m	aterials + 1	0% of other	annual costs									
31	l otal, million\$	6		-t-\:10°	7.34	8.28	9.73	11.08	12.41	13.83	15.15			
32	Total investment mil	i annuai	variable cos	sis), milā	206.67	21.30	25.40	29.35	282.86	300.92	41.00			
24	Unit Cost of Batte	ny Pac	L ¢		200.07	223.20	244.11	204.33	202.00	300.32	517.05			
36	Variable Cost	iyiac	π, ψ											
36	Materials and Purchase	ed Items												
37	Cell materials	, a normo			556	711	940	1,168	1,395	1,621	1,846			
38	Cell purchased Item	IS			73	76	79	82	85	87	90			
39	Module				276	284	292	300	309	317	325			
40	Battery pack				117	120	141	147	153	180	187			
41	Total				1,022	1,191	1,452	1,697	1,941	2,205	2,448			
42	Direct Labor at	18	\$/hour		20	22	20	12	47	E 4	E 4			
43	Cell assembly	ig			30	3/	3/	43	47	3/	34			
45	Formation cycling t	esting a	nd sealing		22	22	22	22	22	22	22			
46	Module and battery	assemb	ly		20	20	20	20	20	20	20			
47	Cell and materials re	ejection	and recyclin	ng	9	9	9	9	9	9	9			
48	Receiving and shipp	ing			5	7	7	8	9	10	10			
49	Control laboratory				4	4	5	6	6	7	7			ļ
50	I otal Variable Overhand				124	128	135	142	147	152	15/			
52	Total Variable Cost				1 245	1 / 2/	1 600	1 956	2 212	2 /86	2 739			
53	Fixed Expenses				1,240	1,424	1,033	1,000	۲,۷۱۷	2,400	2,133			<u> </u>
54	General, Sales, Admini	stration			117	124	133	141	148	155	161			
55	Research and Developn	nent			98	105	114	122	129	136	142			
56	Depreciation				245	263	285	304	322	339	355			
57	Total Fixed Expenses				461	492	531	567	599	630	659			
50	Total unit cost per pack	not incl	uding wars	ntv S	103	2 029	2 252	132	2 052	3 366	3 555			
00	Summany of Unit (Coete	¢	πy, φ	1,005	2,020	2,303	2,000	2,332	3,200	3,550			
61	Materials	00313,	v		973	711	010	1 169	1 305	1 621	1 8/6			
62	Purchased Items				466	480	512	530	547	584	602			<u> </u>
63	Direct Labor				124	128	135	142	147	152	157			
64	Variable Overhead				99	104	111	117	123	129	134			
65	General, Sales, Admini	stration			117	124	133	141	148	155	161			
66	Research and Developn	nent			98	105	114	122	129	136	142			
67	Depreciation				245	263	285	304	322	339	355			
60	Marranty (includes hat	on nor	(c) orby)		103	112	122	132	141	150	159			
70	Price to OEM for batter	v pack	s only		1 911	2 141	2 484	2 80/	3 118	3 449	3 756			
71	Pack integration (BMS	& Disco	<pre>mnects) \$/m</pre>	ack	395	395	395	395	395	395	395			
72	Total battery cost to OE	EM, \$			2,306	2,536	2,879	3,199	3,513	3,844	4,151			
73	Additions to AC for ther	rmal mar	nagement, \$;	200	200	200	200	200	160	160			
74	Total cost to OEM for c	omplete	system, \$		2,506	2,736	3,079	3,399	3,713	4,004	4,311			
75	Price to OEM for modul	les for o	ne pack, \$	0/	1,761	1,989	2,309	2,622	2,929	3,232	3,532			
76	Error bars on price to O	VEIVI for b	battery pack	(, %)	40	07	20	24	20	22	24			
78	Potential positive en	rror %			10	27	29	31 17	JZ 18	33	34			<u> </u>
79	Cell capacity inadequ	uate (if)	X)				10		10	10	10			
00			-											

Figure 6.7	Summary	of Results	worksheet
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6.3 Battery Design Format Requirements

As the battery design is based off an assumed format (Section 2), certain design requirements are necessary to ensure the modeled battery is physically realistic. The dimensions of the calculated battery pack should be examined. Some final designs may benefit from changing the cell aspect ratio, H/W, to fit the end-use application. One example would be, to reduce the height of the battery pack by increasing the cell H/W ratio. Also, for a set number of cells in the pack, changing the number of rows of modules, allows for adjustment of the pack dimensions.

6.4 Troubleshooting and General Advice

The spreadsheet iterates to find the solution and this sometimes causes error messages to appear after an entry is changed. These errors can usually be removed by first correcting any erroneous entries (non-numeric, two decimal points, etc.). Then the cells may be reset to default values by entering a "0" (i.e. zero) in the restart cell, F188 in Figure 6.6. Finally, entering a "1" in F188 restarts the iteration process leading to a successfully converged answer.

At some point, a user will ask the model to design a battery that is outside the bounds of what is allowable for the selected cell chemistry. The most common error is when too large of a P/E ratio is requested. Two different physical limitations are approached with increasing P/E ratio. First, the electrode thickness is shrinking. At some point, the value will become unrealistic and eventually approach 0 crashing the calculation. At the same time, the C-rate for the active material is approaching the limiting C-rate defined in the Chem worksheet. As this value is approached, the ASI will increase to larger and larger values, which thus demands smaller and smaller electrode thicknesses. Eventually, the calculation will crash.

A warning has been added to BatPaC in row 72 of the Battery Design worksheet. A red "X" will appear if the capacity is too low to meet the power required by the user. The warning is triggered if the ASI for power is calculated to be more than 50% the default level (see cell E45 on Chem worksheet). Common sense approaches to resolve these issues are to use lower designed power or higher designed energy. The C-rate and electrode thickness are easily viewed in the model output. These are found on the Battery Design worksheet in row 139 for the C-rate and rows 94 and 95 for the electrode thickness. Therefore, the user may try designs of increasing P/E ratios and watch to see how the electrode thickness and C-rate is changing. Different cell chemistries will have different sensitivities to the P/E ratio depending on the defined limiting C-rate, $r_{C,lim}$, and calculated ASI for power. What is possible with the LMO-G system will not always be possible with the NCA-G system. P/E ratios that satisfy the expression in Eq. 6.1 generally result in successful battery designs. Higher P/E ratios are allowable in some situations. Note that selecting the microHEV design doubles the allowable C-rate since only two second pulses are used. The limiting C-rate, $r_{C,lim}$, may be found in cell E44 on the Chem worksheet and is carried over to row 77 in the Battery Design worksheet.

$$\frac{P}{E} < \frac{r_{C,\text{lim}}}{1.35} \tag{6.1}$$

6.5 Suggested Number of Cells, Modules, and Performance Inputs

Table 6.1 presents some suggestions for the required inputs into the design model that might change depending on the type of vehicle battery being designed. These values are only suggestions, but tend to be similar to practices used today or projected to be used in the future by industry. If the calculated cell capacity is higher than 60 Ah, the user should consider the inclusion of parallel cells as an additional parameter to examine. The default energy usage rate in BatPaC is 300 Wh/mi. This rate may be used to size the energy requirement based on a desired electric range for the vehicle by specifying the distance in row 164 on the Battery Design worksheet.

battery type	modules/battery	OCV @ 50% SOC	Power (kW)	Energy (kWh)
HEV-25	1 - 4	40 - 200	25	0.6 - 1.5
HEV-HP	2 - 4	160 - 260	25 - 80	1 - 2
PHEV	4 - 6	290 - 360	40 - 160	4 - 30
EV	4 - 6	290 - 360	80 - 160	20 - 200

Table 6.1 General suggestions for range of input parameters that change with battery type

6.6 Entering a New Material Couple

The user of the model may wish to examine an electrochemical couple that is not included as one of the options available in the model. We list below a brief explanation on how to properly enter new materials into BatPaC. Various properties may be calculated, found in literature or measured in the laboratory. The self-consistency of the data used is very important.

Experimentally measured values required:

- 1. Half cell formation cycling data from positive and negative electrode
- 2. Half cell cycling data from positive and negative electrode at C/3 rate
- 3. Full cell open-circuit voltage measurement at 50 % and 20 % SOC
- 4. Full cell ASI measurement for 5C pulse at 50 % and 20 % SOC
- 5. Full cell ASI at 50% SOC during a C/3 discharge
- 6. Electrode void fractions, active material densities, electrode component weight percent
- 7. Estimated interfacial area from surface area (preferred) or particle size measurements

The ASI calculation includes some additional parameters that become important as the designed P/E ratio increases above 10 h^{-1} or the electrode thicknesses decrease below 30 microns. The user is referred to section 3.4 and the supporting manuscript from Gallagher *et al.*²⁰ for the parameter estimation process. An exchange current of 0.15 mA/cm², normalized to the surface area calculated using the BET method from nitrogen absorption experiments, is used in the model in row 77 in the Battery Design worksheet. While the exact value of the exchange current will vary from the material to material, the general behavior of the ASI will be preserved with this assumption. If lower P/E ratio designs are desired, the exact valuation of the exchange current, interfacial area, and limiting C-rate are less important. However, the experimental ASI measurement should then come from a cell with similar P/E ratio (electrode loading). Electrode thicknesses 40 microns or larger should be used to minimize the contribution of interfacial

impedance to the ASI measurement. Otherwise the "ASI correction factor" may not accurately remove the interfacial component. A reasonable approach for a first approximation of the ASI parameters may be to select the same values for a similar material. For example, if the new material is based on nano-sized primary particles, then the parameters for LFP or LTO may be close enough. The ASI does depend on a large number of factors and a full determination of the parameters is important to capture all of the physical behavior of the cell couple. If the desire of user is to reproduce an existing cell, then many of the parameters may be estimated from the electrochemical characterization of the full cell data. These electrochemical results typically require a teardown of the cell to measure the area of the electrodes and their loadings/thicknesses.

The available lithium for cycling in a full cell configuration may be calculated from half cell measurements. The calculation method may be found in the Capacity Calculator worksheet in the spreadsheet model and is detailed below. Alternatively, the reversible capacity of the full cell in the experiment may be normalized to the mass of the positive and negative electrodes while carefully accounting for the negative to positive capacity ratio. The first cycle efficiency of a positive or negative electrode based half-cell may be defined as the ratio of the first discharge capacity divided by the first charge capacity, Equation 6.2. We have assumed the first discharge capacity is equivalent to the reversible capacity when measured against lithium foil (half-cell arrangement).

$$\eta_j^{1st} = \frac{Q_j^{rev}}{Q_j^{1st}} \tag{6.2}$$

The quantity of lithium consumed from the positive electrode in the negative electrode SEI, Q^{sei} , may be calculated from Equation 6.3. Here, [N/P] is the negative to positive capacity ratio.

$$Q^{sei} = \left[N/P\right] \frac{Q_P^{rev}}{Q_N^{rev}} \left(Q_N^{1st} - Q_N^{rev}\right) = \frac{m_N}{m_P} Q_N^{irrev}$$
(6.3)

After one full cycle, the remaining lithium in the positive electrode available for cycling, Q_P^{act} , in a full-cell configuration (positive electrode versus non-prelithiated negative electrode) may be calculated by choosing the minimum value determined in Equation 6.4 below. Here we see the possibility that the positive electrode is unable to accept the full amount of lithium released during the first charge cycle. This so called "irreversible capacity" of the positive electrode results in lithium residing in the negative electrode. While this excess lithium may require additional negative electrode capacity, it also provides some beneficial aspects to cycle and calendar life.⁵⁰ We have chosen to set the [N/P] = 1.25 for layered oxides positive electrodes (NCA, NMC441, NMC333) due to their tendency to have a lower first cycle efficiency ~ 88%. The lithium manganese spinel and lithium iron phosphate cells have a high first cycle efficiency and thus we selected a [N/P] = 1.20. For positive electrodes with a high first cycle efficiency, the reversible capacity of the cell is reduced by the lithium consumed in the graphite electrode SEI during the formation cycle. Conversely, the lithium titanate spinel negative electrode does not form an SEI and is significantly safer than the graphite electrode as discussed in Chapter 5. Thefore the [N/P] ratio is set to 1.1 for the cells based on lithium titanate spinel.

$$Q_P^{act} = \mathbf{M} \operatorname{IN}\left\{ Q_P^{rev} \left(1 + \left(\frac{1 - \eta_P^{1st}}{\eta_P^{1st}} \right) - \left[\mathbf{N} / \mathbf{P} \left(\frac{1 - \eta_N^{1st}}{\eta_N^{1st}} \right) \right], Q_P^{rev} \right\}$$
(6.4)

6.7 Entering Parallel Cell, Module, or Pack Configurations

The provision for specifying multiple modules and multiple packs was added to the worksheet "Battery Design" and the equations throughout the spreadsheet were altered as necessary. To implement parallel cells, the user must designate the number of cells in parallel and increase the number of cells in a module accordingly. The model program will automatically select a cell thickness of 6 mm, which is half the thickness selected by the model for EV cells if all are connected in series. Because of the larger number of cells, the price of a battery with parallel connected cells is higher than that for the battery of all series connected cells. To select a parallel module configuration, the user of the model must increase the number of modules and select the desired number of modules in parallel. The cost of the parallel modules is more than that of an equivalent battery with parallel cells because of costs for extra module hardware especially for the extra circuits for SOC regulation.

To a parallel battery pack setup, the user must first configure the battery packs, remembering that each pack provided only a part of the energy, and then select the number of packs and designate whether they are in parallel or series connection. The user must also determine the capacity of the pack at the bottom of the "Battery Design" worksheet. The user makes this determination by selecting one of the following: (1) the pack capacity (note: not the cell or total battery system capacity), (2) the pack energy, or (3) the range of the vehicle. For Table 1, we selected a 100 mile range for all five sample battery systems. The battery system with series-connected packs has twice the voltage and half the system capacity than for a parallel pack configuration.

7. Illustrated Results

The BatPaC model may be used to study the effects of battery parameters on the performance and the manufactured cost of the designed battery packs. A few examples are given below for the effects of various parameters on battery pack volume, weight and cost.

7.1 Number of Cells in Series

For a set battery pack power, the number of cells in the pack has substantial effects on the price of the pack, the pack voltage and the maximum current. These effects are illustrated (Figure 7.1) for LMO-Gr PHEV20 batteries (providing 20-mile electric range) with 60-kW power at a [V/U] = 0.8. The total battery cost to the OEM increases by 15% in changing the number of series-connected cells in the pack from 48 to 96. The integrated cost includes battery management system with disconnects in addition to battery price. The change in the maximum current, resulting from differing pack voltages, would also affect the cost of the motor and the electronic converter and controller, but in the opposite direction. As a result of these offsetting effects on the total cost of the electric drivetrain, a study is required to determine the optimum current at rated power as a function of the total battery pack power and other parameters.



Figure 7.1 The effect of the number of series-connected cells for LMO-Gr, 60-kW, PHEV20 packs with 8 kWh total energy (70% useable).

Current PHEV and EV battery technology uses battery packs containing 80-96 series connected cells. However, these series connections are often composed of parallel cell groups. For instance, the battery used in the first production model of the Chevrolet Volt is in a 3P-96S configuration.²¹ Three low capacity cells are connected in parallel forming a parallel cell group. Then 96 parallel cell groups are connected in series. The cost savings from moving to larger format cells with only series connections is discussed later in this section.

7.2 Cathode Materials

Lithium-ion batteries for high mileage PHEVs and EVs do not require a high P/E ratio or low ASI to meet their goals. The most important material properties for performance are high specific capacity (mAh/g), high cell voltage, and high electrode density. Graphite electrode is near universal in commercial cells as a negative electrode, although not all graphite is the same. Here we consider changes in the positive electrode that result in significantly different calculated batteries. To compare the performance of EV battery packs made from various Li-ion chemistries, we designed the packs to provide 150 kW at 360 V (20% SOC) for a [V/U] = 0.8. Each pack consisted of six modules containing 16, 16, and 18 cells for the cell chemistries LMO-Gr, NMC441-Gr, and LFP-Gr, respectively. This calculation assumes that large capacity cells may be reliably produced. Moving to a parallel connection of smaller capacity cells would result in higher cost as discussed later in the section.

The NMC441-Gr system has excellent energy density and low cost (Fig. 7.2 and 7.3). The LMO-Gr system is less energy dense than the NMC441-Gr couple, but equivalent in calculated price to the OEM. The LFP-Gr system results in a battery that is larger and more expensive than the other two chemistries. The mass-specific cathode raw material prices are 26, 20, and 10 \$/kg for NMC441, LFP and LMO respectively. The differences in cathode materials costs do not directly translate to the end cost of the battery. The performance (exhibited by specific capacity and voltage) affect the quantity of both active and inactive material required. The NMC441 material achieves 175 mAh/g at a good cell voltage and is representative of an advanced, although close to commercialization, layered oxide cathode.⁵¹ The combination in voltage and capacity results in a superior energy density compared to the other cathodes. The LMO cathode has similar cell voltage to NMC441 and low raw material cost but also a low specific capacity of 100 mAh/g. This low capacity results in a positive electrode loading limited by the maximum achievable electrode thickness ~100 microns. The LFP electrode has moderate capacity, 150 mAh/g, and raw material cost, but exhibits a lower cell voltage and electrode density. These poor performance characteristics result in a low energy density battery with a high price.

7.3 Parallel-Connected Cell Groups and Electrode Thickness Limits

BatPaC also allows the user to create parallel cell groups and to set a maximum electrode thickness. The effect these two unique design factors have on battery price are illustrated below in Figure 7.4 for the LMO-Gr and NMC441-Gr systems. In this illustration, the PHEV battery pack design parameters are 100 kW of power at a [V/U] = 0.8 and 17 kWh of total energy. The nominal battery pack voltage (OCV at 50% SOC) is around 360 V from 96 cell groups connected in series. The number of cells in the parallel cell group is varied from a single cell (no parallel connections) to four. Two maximum electrode thicknesses of 100 and 200 microns are shown for the LMO-Gr chemistry. In contrast to the NMC441-Gr, the LMO-Gr chemistry benefits from allowing larger electrode thicknesses. The thickness of the positive electrode is limiting the LMO-Gr chemistry while the thickness of the negative electrode limits the NMC441-Gr cells.



Figure 7.2 Mass and volume of electric vehicle battery packs with lithium iron phosphate (LFP), lithium manganese-spinel (LMO) and lithium nickel-manganese-cobalt oxide (NMC441) positive electrodes versus graphite designed to deliver 150 kW of power at 360 V (20% SOC).



Figure 7.3 Total battery cost to OEM for LFP-Gr, LMO-Gr and NMC441-Gr battery packs for same designs as in Fig. 7.2. NMC441-Gr and LMO-Gr result in nearly the same price.

The calculated value for the NMC441-Gr system never exceeds 100 microns for this P/E ratio. The LMO-Gr is the least expensive in all cases. However, the difference between the two chemistries lessens with smaller limiting electrode thickness. The costs will become even closer for lower designed P/E ratios. In general, thicker electrodes reduce the cost of the battery pack by lessening the amount of inactive materials used (separator, current collector, etc). Moving to 200

microns allows for greater savings in the LMO-Gr design but not the NMC441-Gr design. However, a lower P/E ratio design for NMC441-Gr would take advantage of electrode thicknesses greater than 100 microns.

The cell capacity is shown for the NMC441-Gr case limited to 100 microns. While the exact values will change with cell chemistry, they will all be similar. The cell capacity is reduced by one half as a single cell is added in parallel. This approach is commonly used by cell manufacturers and OEMs that cannot reliably produce or successfully operate cells of high capacity for transportation applications. However, this approach also increases the price of the battery pack. In this example, the price is increased by ~ \$500 when an additional string of cells is incorporated in a parallel arrangement.

The model calculations show that the lowest cost battery pack will utilize thick electrodes and large capacity cells. In current practice, these two approaches have yet to be successfully implemented within the entire community. In the challenge of lowering costs, it is useful to point out the largest gains come from the initial advances (e.g. moving from 100 to 200 micron limit). After that point, the benefits are diminishing.

7.4 Manufacturing Scale

The effects of manufacturing scale come into the cost calculation even if the annual number of packs produced is unchanged, but the design is altered (e.g. power is increased). For a fixed design, the effect of changing the scale of operations depends on the fraction of the total price that is made up of materials costs. Unit materials costs change little with scale whereas the costs per pack for labor, capital and plant area may decline substantially with increasing production rates, especially at low production rates, Fig. 7.5.

The lines in the graphs are for the best-fit power relationships through the data with power factors of -0.076, -0.077, -0.147, and -0.211 from the top curve to that at the bottom. The least negative power factor is for the battery pack with the highest fraction of materials cost in the total pack cost. The more negative power factors result from a decreasing contribution of materials cost as a fraction of the total pack cost. These power factors for equations of the cost of a single unit can be converted to factors relating the total annual cost of manufacturing similar to Eq. 5.2 by adding 1.0 to each power factor. Thus the factors become 0.924, 0.923, 0.853, and 0.789. These large factors show only a small to moderate effect of scale. When the power curves are compared to the points in each of the graphs of Fig. 7.5, it is apparent that the scale factors approach one as the scale increases. This is because the model assigns a value of 0.95 for the active materials and 1.0 for the balance of the materials. As the production level increases and the materials costs become a larger fraction of the total price of the battery, the scaling power approaches 1.0 and the effect of scale become very small. Likewise, the effect of scale on battery price is much larger for HEV batteries than for EVs because materials costs constitute a smaller portion of the total cost for HEV batteries. Increasing the production rate for HEV batteries will result in a more dramatic reduction in cost than increasing the production rate for EV batteries.



Figure 7.4 Battery pack cost as a function of number of parallel cells and for different maximum electrode thicknesses. The electrode thicknesses (100 or 200 μ m) represent the limitation not the exact value calculated by the model. The cell capacity is also shown for the NMC441-Gr battery.



Figure 7.5 The effects of manufacturing rate on the price calculated by the model for battery packs of various cell chemistries, power capabilities and vehicle types. The EV and PHEV batteries are composed of 96 cells and the HEV-25 is composed of 48 cells.

8. Statement of Copyright

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Appendix A: BatPaC v1.0 Variation Study

Summary

The following ranges in Table A1 are suggested as an initial approximation to account for variation that will occur in the cost inputs and possible design limitations in BatPaC. The larger uncertainty for LMO in PHEV and EV batteries is due to an interaction with the electrode thickness limitation. Future versions of BatPaC will include error estimation within the model.

Table A1. Suggested confidence bounds as a percentage of the calculated point estimate for a graphite based Li-ion battery using the default inputs in BatPaC.

		Confider	nce Interval
Battery type	Cathodes	lower	upper
HEV	LMO, LFP, NCA, NMC	-10%	10%
PHEV, EV	NMC, NCA	-10%	20%
PHEV, EV	LMO, LFP	-20%	35%

Discussion

The general approach in this variation study is to produce a distribution of cost inputs and analyze a range of battery designs to capture the spread in calculated battery price if the default cost inputs or design parameters are in error. The eight largest cost contributions comprising the battery price to the OEM, shown in Table A2, were evaluated to determine the range of possible values and the likely form of the distribution, normal or log-normal. For normal distributions, the standard deviation was approximated as ¹/₄ of the range of values. The parameters for log-normal distributions were hand-fit to set the minimum and maximum at the outer range of the 95 % confidence interval. Equations A1 and A2 were used to create the distribution for each cost input, C_{ij} , from independent sets of 3000 normally distributed random numbers, r_{ij} . Here, C_i is the mean and σ_i is the standard deviation for the normal distribution of cost input *i*. Additionally, μ_i is the location parameter and α_i is the scale parameter for the log-normal distribution. Each collection of cost inputs was used to produce an end battery price to the OEM. The battery designs used in this study are detailed in Table A3.

$$C_{ij} = C_{i,mean} + r_{ij}\sigma_i$$
 normal (A1)

$$C_{ij} = exp(\mu_{i,mean} + r_{ij}\alpha_i)$$
 log-normal (A2)

The variation in calculated price solely due to cost input estimates results in a near-normal distribution of values. The results may be adequately summarized by the calculated mean and standard deviation. Only the cases of the default design limitations are shown in Table A4 (100 micron limit, 1 cell in cell group). A value of \pm 10 % would adequately capture the 95% confidence interval of the calculated price distributions for all battery chemistries and battery types.

Item	low	medium	high	distribution	mean	std dev	location	scale
Separator	0.75	2	4	log-normal			0.75	0.3
Copper foil	1	1.8	2.6	normal	1.8	0.4		
Electrolyte	18	21.6	25.2	normal	21.6	1.8		
Graphite anode	12	19	25	normal	19	3.25		
LMO cathode	8	10	16	log-normal			2.35	0.15
NMC cathode	22	29	34	normal	29	3		
NCA cathode	22	37	42	normal	37	5		
LFP cathode	10	20	30	normal	20	5		
SOC controller	1.5	2.5	3.5	normal	2.5	0.5		
Coating capital	5	8	12	normal	8	1.75		
Formation capital	20	30	50	log-normal			3.45	0.2

Table A2. Cost input ranges and distributions used in variation study

Table A3. Battery designs considered in variation study

Battery type	HEV-HP	PHEV25	EV100
Power (kW)	40	65	115
Energy (kWh)	1.5	10.7	30
Range @ 300 Wh/mi	1	25	80
Number of cells	72	96	96
Number of modules	4	6	6
Cells in parallel group	1	1	1, 2
Electrode thickness limit (micron)	100	50, 100, 150	70, 100, 200
Cathode chemistries	LMO, LFP	LMO, NMC441, NCA	LMO, NMC441, NCA

Table A4. Distribution in calculated price from cost input variation

battery	cathode	mean	std deviation	2σ/mean
HEV	LMO	1268	49	0.08
HEV	LFP	1475	65	0.09
PHEV25	LMO	2995	134	0.09
PHEV25	NMC441	3087	128	0.08
PHEV25	NCA	3325	153	0.09
EV100	LMO	6051	339	0.11
EV100	NMC441	5927	279	0.09
EV100	NCA	6648	364	0.11

Quantifying the error in calculated price due to the relative success or failure of certain design criteria is more difficult. We have chosen to vary the electrode thickness limitation for the PHEV and EV calculations. In addition, the EV battery was also calculated in a 2P-96S configuration (2 cells in parallel) to capture the possibility of smaller capacity cells while maintaining the 100 micron thickness limit. No design changes were analyzed for the HEVs. Treating each of the battery design occurrences as equally probable, the calculated population of each of the chemistries becomes a skewed distribution. Capturing the skewed and sometime bi-modal distribution in a simple statistical fashion is not straightforward. We suggest the use of a distribution parameter, ε , to establish the outer limits of a 95 % confidence interval from the point estimate, P_o , created from the default values. This parameter is used in an empirical fashion in Equation A3 and A4. The factor of 1.05 is used to capture the increase in the mean of the population as compared to a spot estimate using the default inputs. The distribution parameters used in this study are twice the value of the standard deviation divided by the mean.

$$P_{97.5} = 1.05P_o(1+\varepsilon)$$
 (A3)

$$P_{2.5} = 1.05 P_o / (1 + \varepsilon)$$
 (A4)

The NMC441 and NCA cathodes for PHEV and EV data sets may be captured using a distribution parameter of 0.15. The LMO cathode requires a distribution parameter of 0.28. The larger variation of the LMO cathode results from interaction with the limiting electrode thickness. A similar behavior is expected for LFP in energy batteries as this cathode also has a low volumetric capacity. The NMC441 and NCA cathodes have a significantly higher volumetric capacity and thus are less sensitive to this limitation. Figures A1-A3 present the variation of the PHEV and EV trials with the suggested confidence bounds highlighted by red lines and calculated point estimates from the default inputs are represented by the black line.



Figure A1. PHEV25 price distributions for NMC441 / Gr batteries. Red lines denote suggested upper and lower bounds for confidence interval. Black line represents price calculated from default values.



Figure A2. PHEV25 price distributions for LMO / Gr batteries. Red lines denote suggested upper and lower bounds for confidence interval. Black line represents price calculated from default values.



Figure A3. EV100 distributions for LMO / Gr and NMC441 / Gr batteries. Red lines denote suggested upper and lower bounds for confidence interval. Black line represents price calculated from default values.



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