AN EXPERT SYSTEM FOR THE IDENTIFICATION OF SILICON IMPURITIES USING FTIR SPECTROSCOPY

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An Expert System for the Identification of Silicon Impurities Using FTIR Spectroscopy

Ronald E. Perrin and Steven B. Fairchild

An expert-system for the analysis of the infra-red spectra of silicon based electronic materials has been developed. The Fourier Transform Infra-Red Spectroscopy system produces characteristic spectral lines for impurities present in silicon materials under analysis. A MicroVax workstation with a commercially available expert-system shell is attached to the spectrometer. By examining these spectral line patterns the expert-system is able to deduce the identities and concentrations of the unknown impurities. The use of this program greatly reduces the analysis time required for each sample and produces more consistent results.
# TABLE OF CONTENTS

<table>
<thead>
<tr>
<th>SECTION</th>
<th>PAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0 INTRODUCTION</td>
<td>1</td>
</tr>
<tr>
<td>1.1 BACKGROUND</td>
<td>2</td>
</tr>
<tr>
<td>1.2 PROBLEM STATEMENT</td>
<td>4</td>
</tr>
<tr>
<td>1.3 PROJECT OBJECTIVES</td>
<td>5</td>
</tr>
<tr>
<td>2.0 TUTORIAL</td>
<td>9</td>
</tr>
<tr>
<td>2.1 THE REASONING DIMENSION</td>
<td>10</td>
</tr>
<tr>
<td>2.2 THE REPRESENTATION DIMENSION</td>
<td>10</td>
</tr>
<tr>
<td>2.3 EXAMPLE</td>
<td>11</td>
</tr>
<tr>
<td>3.0 KNOWLEDGE BASE DESIGN</td>
<td>17</td>
</tr>
<tr>
<td>3.1 THE OBJECT NETWORK</td>
<td>19</td>
</tr>
<tr>
<td>3.2 THE RULE NETWORK</td>
<td>23</td>
</tr>
<tr>
<td>3.3 PROGRAM FLOW</td>
<td>34</td>
</tr>
<tr>
<td>4.0 SAMPLE RUN</td>
<td>49</td>
</tr>
<tr>
<td>5.0 TESTING THE EXPERT SYSTEM</td>
<td>51</td>
</tr>
<tr>
<td>5.1 DISCUSSION</td>
<td>52</td>
</tr>
<tr>
<td>6.0 FUTURE PLANS</td>
<td>56</td>
</tr>
<tr>
<td>APPENDIX A FORTRAN SUBROUTINES</td>
<td>57</td>
</tr>
<tr>
<td>APPENDIX B LIST OF RULES</td>
<td>76</td>
</tr>
<tr>
<td>REFERENCES</td>
<td>116</td>
</tr>
</tbody>
</table>
# LIST OF ILLUSTRATIONS

<table>
<thead>
<tr>
<th>FIGURE</th>
<th>PAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>11</td>
</tr>
<tr>
<td>The symbolic representation of a class with its related objects and properties.</td>
<td>11</td>
</tr>
<tr>
<td>2</td>
<td>13</td>
</tr>
<tr>
<td>A silicon spectrum showing evidence of the presence of the impurity indium.</td>
<td>13</td>
</tr>
<tr>
<td>3</td>
<td>14</td>
</tr>
<tr>
<td>Symbolic representation of the class INDium_LINES.</td>
<td>14</td>
</tr>
<tr>
<td>4</td>
<td>19</td>
</tr>
<tr>
<td>The class LINES.</td>
<td>19</td>
</tr>
<tr>
<td>5</td>
<td>20</td>
</tr>
<tr>
<td>The class AL_LINES.</td>
<td>20</td>
</tr>
<tr>
<td>6</td>
<td>22</td>
</tr>
<tr>
<td>The class LINES, its properties, subclasses, and objects.</td>
<td>22</td>
</tr>
<tr>
<td>7</td>
<td>24</td>
</tr>
<tr>
<td>The class POSSIBLE_Peaks.</td>
<td>24</td>
</tr>
<tr>
<td>8</td>
<td>28</td>
</tr>
<tr>
<td>The class POSSIBLE_AL_Peaks.</td>
<td>28</td>
</tr>
<tr>
<td>9</td>
<td>36</td>
</tr>
<tr>
<td>The rule START_THINGS_OFF and the branch of the rule network triggered by the hypothesis PEAKS_LOADED.</td>
<td>36</td>
</tr>
<tr>
<td>10</td>
<td>37</td>
</tr>
<tr>
<td>The PEAKS_EXAMINED branch of the rule network.</td>
<td>37</td>
</tr>
<tr>
<td>11</td>
<td>38</td>
</tr>
<tr>
<td>The PEAKS_ANALYZED branch of the rule network.</td>
<td>38</td>
</tr>
<tr>
<td>12</td>
<td>42</td>
</tr>
<tr>
<td>Forward chaining from the hypothesis ANALYZED.</td>
<td>42</td>
</tr>
<tr>
<td>13</td>
<td>44</td>
</tr>
<tr>
<td>The aluminum rules.</td>
<td>44</td>
</tr>
<tr>
<td>14</td>
<td>45</td>
</tr>
<tr>
<td>The boron rules.</td>
<td>45</td>
</tr>
<tr>
<td>FIGURE</td>
<td>PAGE</td>
</tr>
<tr>
<td>--------</td>
<td>------</td>
</tr>
<tr>
<td>15</td>
<td>46</td>
</tr>
<tr>
<td>16</td>
<td>47</td>
</tr>
<tr>
<td>17</td>
<td>48</td>
</tr>
<tr>
<td>18</td>
<td>55</td>
</tr>
</tbody>
</table>

The gallium rules.
The indium rules.
The carbon and oxygen rules.
Possible peak boundaries in a noisy spectrum.
## LIST OF TABLES

<table>
<thead>
<tr>
<th>TABLE</th>
<th>PAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Results of the Al feasibility survey.</td>
</tr>
<tr>
<td>2</td>
<td>All of the peaks that can occur in a silicon spectrum due to the presence of the impurities.</td>
</tr>
<tr>
<td>3</td>
<td>The report generated by the expert system.</td>
</tr>
<tr>
<td>4</td>
<td>Results of the gallium concentration study test.</td>
</tr>
</tbody>
</table>
1.0 INTRODUCTION

The Wright Laboratory, Electromagnetic Materials Division (WL/MLP) conducts research to develop electronic materials of future interest to the Air Force. These materials can include high purity silicon and gallium arsenide. These materials are used as infra-red detectors in space based applications as well as being used as a basis material for high-speed integrated circuits in the next generation of computers. The impurities and defects in these materials determine the operational limitations of the devices constructed from them. One of the optical characterization techniques used to detect these defects is low temperature Fourier Transform Infra-Red spectroscopy (FTIR). The material being examined is cooled to a cryogenic temperature and then excited by an infra-red light source. The absorption and transmittance of infra-red radiation by these impurities and defects produces a unique spectrum. By examining the spectrum, an experienced spectroscopist can determine the presence of each impurity as well as its concentration in the material\textsuperscript{1}. The locations of peaks, the relative peak heights, and the shape of the background in the resulting spectral plot are used to characterize the material's properties. The presence of unwanted impurities in these materials degrades the performance of the end application devices.
1.1 BACKGROUND

Modern research and development laboratories can benefit from applying new automation techniques. The application of these new technologies can result in more accurate, more consistent, and faster data acquisition, thus maximizing personnel resources. WL/MLP decided that an Expert System application to Fourier Transform Infra-Red Spectroscopy (FTIR) would be an optimum project to pursue in terms of AI applications in the modern laboratory. In 1986 and 1987, two separate groups of artificial intelligence (AI) experts from universities and industry surveyed the research being performed at WL/MLP. All current and future areas of research being performed within the WUD 48 and WUD 50 in-house laboratories were examined. The groups were asked to identify laboratory systems which could benefit from the application of artificial intelligence technology and recommend useful hardware and software products. After interviewing the researchers and being briefed on the details of each laboratory's data acquisition and analysis equipment, the AI working groups generated final reports. These reports outlined which research areas would benefit from applying AI technology and produce the greatest pay-off to the Air Force. This pay-off would be realized in terms of higher productivity and more accurate data analysis if automated using AI technology. The results of one survey are listed in Table 1. Note that the areas referenced as FTIR-2 and FTIR-3 show the highest ranking in terms of feasibility and pay-off. We proposed that an expert system that
Table 1. Results of AI feasibility survey.

OVERALL CONCLUSIONS AND RECOMMENDATIONS
The rankings given in the table for the various AI application opportunities are based on technical considerations alone. (1 = highest potential)

<table>
<thead>
<tr>
<th>Opportunity Number and Name</th>
<th>Recommendation</th>
<th>Rank</th>
</tr>
</thead>
<tbody>
<tr>
<td>MBE-1 Utilize Past Experience</td>
<td>pursue/guarded/ build data-base first</td>
<td>3</td>
</tr>
<tr>
<td>MBE-2 Interp. RHEED Oscill.</td>
<td>Do not pursue re-examine later</td>
<td></td>
</tr>
<tr>
<td>MBE-3 Interp. Optical Probes</td>
<td>Do not pursue re-examine later</td>
<td></td>
</tr>
<tr>
<td>MBE-4 Qual. Simul. of Growth</td>
<td>Pursue/guarded/ build data-base first</td>
<td>4</td>
</tr>
<tr>
<td>PVD-1 Correction Analysis</td>
<td>Do not pursue</td>
<td></td>
</tr>
<tr>
<td>PVD-2 Detection Analysis</td>
<td>Do not pursue</td>
<td></td>
</tr>
<tr>
<td>PVD-3 Intell. Front-End to Chem. Abstracts Data-Base</td>
<td>Pursue with constraints</td>
<td>2</td>
</tr>
<tr>
<td>FTIR-1 Initial Set of Experiments</td>
<td>Do not pursue</td>
<td></td>
</tr>
<tr>
<td>FTIR-2 ID Known Centers and Locate Unknown Centers</td>
<td>Pursue</td>
<td>1</td>
</tr>
<tr>
<td>FTIR-3 Categorize and Assess Significant Unknown Centers</td>
<td>Pursue</td>
<td>2</td>
</tr>
<tr>
<td>FTIR-4 Correlate Unknown Centers</td>
<td>Pursue/guarded build data-base first</td>
<td>3</td>
</tr>
<tr>
<td>FTIR-5 Formulate Strategy for Further Experiments</td>
<td>Pursue/guarded/ build data-base first</td>
<td>4</td>
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</tbody>
</table>
could address the areas described in Table 1 as FTIR-2 and 3 be designed. This project would not only benefit the FTIR laboratory but lessons learned would be extremely valuable for future AI projects within WL/MLP.

1.2 PROBLEM STATEMENT

The task of analyzing FTIR spectra is complex and tedious. Simply matching peaks with known impurity wavenumber locations tells the researcher very little. The most common impurities, aluminum, boron, gallium, indium, carbon and oxygen, all occur in the same area of the spectrum (0-2000 wavenumbers). The presence of a singularity, or peak, in the spectrum at a certain wavenumber could be the result of dust present in the sample chamber or it could indicate the presence of an impurity or sample defect. The thickness of the sample as well as the method of mounting it also affects the shape of the spectrum. Sixty-hertz pickup and its harmonics show up in spectra as well as noise spikes that occasionally occur due to power line transients. The researcher must carefully look for peak patterns knowing from experience that many false conclusions could be reached if certain rules of thumb are not followed. It is this well defined set of rules and heuristics which, if incorporated into a knowledge base, could allow a less experienced researcher or aide to accurately characterize samples. This type of knowledge base could be useful for basic research as well as for routine material quality control testing.
Since the area of a selected peak is used to calculate its impurity concentration, the expert system must be capable of defining the baseline of a peak. While this task may seem simple, it must be remembered that peaks can be very weak if the concentrations are small. Peaks can also occur in very noisy areas of the spectrum or may be present where the spectral background is increasing or decreasing. Choosing the baseline of a peak in a noisy area is a subjective choice based on the past experience of the researcher. Often two expert spectroscopists will calculate peak areas differing by 30% or more when examining the same peak in a noisy background. The process of calculating peak areas accurately can be extremely time consuming. Researchers have, in the past, resorted to cutting the peak of interest out of a spectral plot and weighing it on a set of sensitive scales. This weight was compared to the weight of a unit area from the same type of graph paper. Obviously an easier, more consistent method is needed. The application of the proposed expert-system would generate more consistent analyses and make a tedious and exacting task quick and routine.

1.3 PROJECT OBJECTIVES

One of the most difficult tasks in designing this project was deciding what the expert system should actually be able to do. There is always a trade-off between what is needed by the end user and what is feasible to design given the constraints of the expert-system shell, the skills of the designer, and a reasonable completion
There is a saying, "Anything can be made to work if you tinker with it long enough." We wanted to limit the scope of this project to be realistic given our expert system design experience, time constraints, and our other job responsibilities.

It was decided that the expert system, given an FTIR absorption coefficient spectrum, should be able to identify all impurities present, their concentrations in the material, and all the associated impurity lines present. If the spectrum is a photoconductivity spectrum, it should be able to identify all impurities present and their associated line numbers.

To accomplish this, a knowledge base had to be designed that incorporated all the necessary information about how to identify these impurities in a silicon spectrum. The theoretical background for this knowledge base came from two sources. The first source is the knowledge obtained from extensive question and answer sessions with experienced spectroscopists in this particular area of materials analysis. This part is typically referred to as 'knowledge engineering.' The second source of information is technical papers and reports directed in this area of research. Together these sources provide a realistic starting point for constructing a useful expert system. Table 2 is taken from such a paper and lists all the lines (i.e., peaks) by wavenumber that can occur in a silicon spectrum owing to the impurities of interest.

There should be minimal user interaction. The user could be asked if they agree with interim results; if not, the user has the authority to modify these, but the final conclusions are determined by the expert system. A graph of the spectrum being examined should be
Table 2. All of the peaks that can occur in a silicon spectrum due to the presence of the impurities.

<table>
<thead>
<tr>
<th>Line</th>
<th>BORON</th>
<th>ALUMINUM</th>
<th>GALLIUM</th>
<th>INDIUM</th>
<th>THALLIUM</th>
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<td>442.73</td>
<td>469.75</td>
<td>1145.41</td>
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<td>1175.82</td>
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<td>309.32</td>
<td>516.52</td>
<td>548.23</td>
<td>1208.00</td>
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<tr>
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<td>299</td>
<td>454</td>
<td>909</td>
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</table>
displayed with all peaks marked by the expert system. If any peaks of interest have been missed, the user should be able to add these with the mouse and cursor. The user should not be allowed to delete peaks that the system has marked; it is up to the expert system to conclude whether marked peaks are significant. All impurities identified must be present in the patterns that experienced spectroscopists recognize as being necessary to insure a high degree of confidence. Noise and spurious peaks present in the spectrum should be ignored. Indeterminate peaks should be noted and classified as unknown. Finally, a report containing this information should be prepared in a concise format and be displayed and printed upon conclusion of the analysis.
2.0 TUTORIAL

The purpose of an AI knowledge based system is to transform the line of thought of a human expert into a set of formalized symbolic operations so as to reproduce his or her expertise. Knowledge design is the process of transferring this formulated knowledge from humans to the machine. Once the computer has this set of rules formulated from human logic, it can perform deductive or inductive reasoning and arrive at the same conclusion as the human expert when given the appropriate data.

The methodology involved in knowledge engineering involves two closely related critical dimensions: the reasoning dimension and the representation dimension. The reasoning dimension is composed of the rules that constitute the logic the expert system uses to perform reasoning. The representation dimension is composed of objects and classes, which can be thought of as symbolic representations of the data that the rules reason upon. Therefore, objects and classes are complimentary to rules.

The reasoning dimension and the representation dimension are closely interrelated to provide an efficient network. They allow ease in knowledge integration, which is the reduction of thought mechanisms to more understandable concepts. Once knowledge integration has been accomplished, powerful symbolic computations can be performed which allow automation of the problem solving process.
2.1 THE REASONING DIMENSION

To represent reasoning, an expert system uses rules. Rules are symbolic structures which express deductive or evocative progressions in a reasoning path. By deductive we mean that a rule can be used to verify conditions in another rule. By evocative, we mean that a rule can trigger the activation or the evaluation of other rules.3

A rule is a chunk of knowledge representing a situation and its immediate consequences. The format of a rule is a symbolic structure of the type

if ... then ... and do...

where if is followed by a set of conditions, then by a hypothesis or goal which becomes TRUE when the conditions are met, and do by a set of actions to be undertaken as a result of a positive evaluation of the conditions of the rule.3

2.2 THE REPRESENTATION DIMENSION

The representation dimension is composed of a set of interrelated objects which represent the data or events that the rules act upon. Since the rules reason on a representation of the problem, the reasoning and representation dimension are inherently related.3

The representation of knowledge is accomplished with three symbolic tools: objects, properties, and classes. Various
combinations of these three components provide an efficient method of representing the problem upon which the rules act.\textsuperscript{3}

An object is an elementary unit of description. Everything is an object. A property is a characteristic of an object. A class is a collection of objects that share properties. As illustrated in Figure 1, objects are illustrated by triangles, properties by squares, and classes by circles.\textsuperscript{3}

![Diagram](image)

Figure 1. The symbolic representation of a class with its related objects and properties.\textsuperscript{3}

2.3 EXAMPLE

This relationship between classes, objects, and properties and their connection to rules can be seen in the following example, which introduces the concept of knowledge base design to spectroscopic analysis.
A spectroscopist can examine the optical spectrum of a material to determine its composition. For example, pure silicon can always be identified by its specific characteristic spectrum. If a certain spectrum of silicon contains peaks that would not show up in a spectrum of pure silicon, then there are impurities present in the silicon sample from which the spectrum was generated. From previous experimental knowledge we know that various impurities identify themselves by the specific wavenumbers at which these peaks appear.

For example, if the impurity indium is present in a silicon sample, it can manifest itself by causing as many as 20 additional peaks to appear in the spectrum, three of which can be used to confirm the presence of indium. These peaks will be referred to as Indium_Lines. The confirming peaks are Indium_Line_1, Indium_Line_2, and Indium_Line_4. All three of these lines must be present for indium to exist. Figure 2 shows an example of a peak pattern in a silicon spectrum that identifies the presence of indium.

Figure 3 illustrates the object structure. Indium_Lines is a class, and Indium_Line_1, 2 & 4 are the objects. As previously stated, a class is a group of objects that have the same properties. Since each object Indium_Line_n will be identified by a specific wavenumber, each object will be given the property WN, for wavenumber. Each object will also be given the property EXIST, which will be used to identify whether or not that particular peak is present.
Figure 2. A silicon spectrum showing evidence of the presence of indium.
Figure 3. Symbolic representation of the class Indium_Lines.

Rules must operate on data, and data in the object network is represented by *slots.* A slot is the value that is held in the property of an object:

\[ \text{object.property} = \text{slot} \]

For this object network, a slot would be Indium_Line_1.EXIST and this slot (datum) would have a boolean value. Another slot would be Indium_line_4.WN, and it would have a numeric value. For example, the wavenumber for the third indium line is 1216, so the value held in the slot Indium_line_4.WN is 1216.

Rules operate on the objects, which are a symbolic representation of the problem. A rule can be designed that will confirm whether or not indium is present in the silicon spectrum. This rule attempts to confirm the presence of the confirmation pattern.
if Indium_Line_1.EXIST is true
and Indium_Line_2.EXIST is true
and Indium_Line_4.EXIST is true
then Indium_Present is confirmed

In this rule, there are three *conditions* which lead to the evaluation of a *hypothesis*. The conditions are the tests for the presence of the three indium lines, and the hypothesis is the attempt to confirm the presence of indium. All three conditions must be true in order for the hypothesis to be confirmed, i.e., all three indium lines must be present in order for the presence of indium to be confirmed.

This rule operates on the slots that were set up in the object network. It will evaluate the slots Indium_line_n.EXIST. (Assume that a previous rule has determined the values for Indium_line_n.WN.) The values of these slots will determine whether or not the hypothesis is confirmed. The name of the hypothesis is Indium_Present, which has a Boolean value. If all the data values (i.e., Indium_line_n.EXIST) that occur in the *conditions* evaluate to true, then the hypothesis Indium_Present will evaluate to true. If any of the datum evaluate to false, the hypothesis Indium_Present will evaluate to false.

Therefore, if the pattern of the three indium lines is found, this rule will confirm that indium is indeed present. If any one of the three lines does not exist, then the rule confirms that indium is not present, since the whole pattern must be present to confirm the presence of indium.
This example has introduced the concept of knowledge base design. The most important part of knowledge base design is the inherent relationship between rules and the object network. As illustrated, the object network is composed of classes, objects, and properties, which are a symbolic representation of the problem. Rules can be related to the object network through their ability to evaluate data. Data are represented by a slot, which is the value stored in a property attached to an object.
3.0 KNOWLEDGE BASE DESIGN

As previously stated, the critical task of this project is to design the knowledge base so that the expert system will have the ability to identify all impurities of interest that can be present in a silicon sample. These impurities are aluminum, boron, gallium, indium, carbon, and oxygen. The presence of these impurities can be confirmed by identifying the variations they cause in a silicon absorption spectrum. This confirmation process involved three critical steps: (1) defining the wavenumber range where the peaks occur for each impurity, (2) identifying the patterns that confirm their existence, (3) and classifying all additional lines associated with that impurity which are not contained within the identifying pattern.

For the first step, the wavenumber ranges of the various impurities must be identified. From previous experimental knowledge, it is known that each possible impurity causes peaks to occur in a well defined region. For example, if aluminum is present in a silicon sample, peaks will appear in the spectrum in a range from 337 to 912 wavenumbers. As many as 20 different peaks can show up in this range. The problem lies in determining which peaks present in this range are actually due to the presence of aluminum and which ones are caused by extraneous noise.

This leads to the second step, identifying the patterns of each impurity to confirm its presence. Each of the 20 aluminum peaks that may appear can be distinctly defined by a particular wavenumber, which we will call aluminum_line_n. If aluminum is
present, it is known from previous experimental knowledge that the pattern of aluminum_lines 1, 2, and 8 must appear in the spectrum. If strong identifiable peaks appear around 442 wavenumbers (aluminum_line_1), 471 wavenumbers (aluminum_line_2), and 543 wavenumbers (aluminum_line_8), then you can positively conclude that aluminum is present.

After this pattern matching has accomplished the identification of the impurity, the next step is to identify all additional peaks that are associated with that impurity. For aluminum, there may be as many as 17 peaks, since only 3 out of a possible 20 were needed to identify the pattern. These peaks must be identified by their wavenumbers and classified as aluminum_line_n. The important point to remember is that there may be peaks that show up at the aluminum_line_n wavenumbers, that are not due to the presence of aluminum. If the pattern of aluminum_lines 1, 2, and 8 has not been previously confirmed, the presence of these peaks must be caused by some extraneous noise. Any peaks that show up at the aluminum_line_n wavenumbers in the absence of the confirmation of the aluminum pattern would be considered unidentified.

In summary, the goal of this effort is to accomplish the impurity analysis for the six possible impurities of interest that can occur in silicon. These are aluminum, boron, gallium, indium, carbon and oxygen. The identification process is the same for each impurity. First, the wavenumber range where they will cause peaks to occur must be defined. Next, the patterns of peaks that confirm their existence must be identified. And finally, all additional peaks
that are caused by the impurity which are not contained in the identifying pattern must be classified.

3.1 THE OBJECT NETWORK

As stated in the tutorial section, the object network represents the representation dimension. The representation is composed of classes, objects, and properties which can be thought of as symbolic representations of the data that the rules reason upon. The object network designed for this situation must therefore use classes, objects, and properties to represent the task of using spectral analysis to identify impurities in silicon.

The peaks that appear in a silicon spectrum that are due to the presence of impurities are classified as lines. Therefore a class will be used called LINES. It will have six subclasses for the six impurities that may be present, aluminum, boron, gallium, indium, carbon, and oxygen (Figure 4). These subclasses will be called AL_LINES, BO_LINES, GA_LINES, IN_LINES, CA_LINES, and OX_LINES. Classes are symbolically represented by circles.

Figure 4. The class LINES
Each subclass will contain objects to represent all of the lines whose existence may be due to the presence of that impurity. For example, the subclass AL_LINES (Figure 5) will contain the objects AL_LINE_1, AL_LINE_2, etc. Each one of these objects has two properties, EXIST, and WN (wavenumber). EXIST will have a Boolean value and WN will have a numeric value. The slot AL_LINE_1.EXIST will be set to TRUE if that particular line exists. If the line exists, the slot AL_LINE_1.WN will be given the value of the wavenumber where the peak was found. Objects are symbolically represented by triangles and properties by squares.

Figure 5. The class AL_LINES.
In the previous diagram, the properties EXIST and WN are shown attached to the class AL_LINES, rather than to the individual objects. This illustrates the principle of downward inheritance. If properties are initially attached to a class, then all the objects within that class will inherit these properties. This is very efficient for the situation where all of the objects in a class are similar items with the same characteristics. The properties EXIST and WN don't have to be individually attached to each of the 20 aluminum lines. As long as they are attached to the class, these properties will be inherited by all 20 objects which represent the aluminum lines.

Figure 6 shows the full extent of the class LINES. It is shown with all of its subclasses, and all of the objects that are contained within each subclass. As shown, downward inheritance also occurs from class to subclass. Initially, the properties EXIST and WN are attached to the class LINES. They are then inherited down by the subclasses AL_LINES, BO_LINES, etc., and from there they are inherited down to each individual object of all of the subclasses.

The objects illustrated represent all of the lines that the expert system will attempt to identify. Aluminum, boron, gallium, and indium are responsible for the majority of the impurity induced peaks that can occur in a silicon spectrum. Carbon and oxygen are responsible for causing a peak to occur at only one wavenumber each if they are present.
Figure 6. The class LINES, its properties, subclasses, and objects.
The presence of impurities in a sample of silicon may cause as many as 84 different peaks to appear in its spectrum. The class LINES contains six subclasses which contain all of the lines that are potentially caused by these impurities.

3.2 THE RULE NETWORK

The rule network represents the reasoning dimension. The reasoning dimension is composed of the rules that constitute the logic the expert system uses to perform reasoning. Rules were designed to categorize the peaks according to the wavenumber ranges of the impurities of interest, and also to identify their other lines if their patterns were recognized.

First, the expert system must obtain a list of the peaks that appeared on the silicon spectrum. After the spectrum has been generated on the FTIR spectrometer, a program is run which locates the peaks in the spectrum, and lists them in a table by wavenumber and amplitude. This peak table has the following format:

<table>
<thead>
<tr>
<th>name</th>
<th>wavenumber</th>
<th>amplitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak_1</td>
<td>432.56</td>
<td>0.654</td>
</tr>
<tr>
<td>Peak_2</td>
<td>458.78</td>
<td>0.421</td>
</tr>
<tr>
<td>Peak_3</td>
<td>543.89</td>
<td>0.784</td>
</tr>
</tbody>
</table>

If \( n \) peaks were found in the spectrum, \( n \) peaks will be listed in this peak table in ascending order by wavenumber. Peak_1 would have the lowest wavenumber and Peak_\( n \) the highest.
This peak table is read into the knowledge base. A class is created called POSSIBLE_PEAKS, and each peak is an object of this class (Figure 7). The name of the objects will be PEAK_1 through PEAK_n, and they will have the properties WAVENUMBER and AMPLITUDE. Using the previous table as an example, the slot PEAK_1.WAVENUMBER would contain the value 432.56, and the slot PEAK_1.AMPLITUDE would contain the value 0.654. With this symbolic representation of the initial peak data, the rules can now begin the reasoning process.

![Figure 7. The class POSSIBLE_PEAKS.](image)

The first task of the rule network is to categorize the peaks in terms of what impurity they might possibly represent. Since the wavelength range for aluminum peaks is 337 to 912 wavenumbers, a rule must be written that finds all of the peaks in the class
POSSIBLE_PEAKS that fall within this wavelength range. The name of this is PICK_POSSIBLE_ALUMINUM_PEAKS.

This rule will utilize a technique called pattern matching, which permits the creation of a list of objects that satisfy a certain set of conditions. First, the rule will sort through all the peaks in the class POSSIBLE_PEAKS and keep the ones whose wavenumber is greater than 337. Then it will sort through that list again and keep all of the peaks whose wavenumber is less than 912. Then it will take this final list and use it to create a new class called POSSIBLE_AL_PEAKS. This new class will contain all of the peaks whose peak location falls within the range of 337 to 912 wavenumbers. The rule looks as follows:

Name: PICK_POSSIBLE_ALUMINUM_PEAKS

Conditions:

> <POSSIBLE_PEAKS>.WAVENUMBER 337
< <POSSIBLE_PEAKS>.WAVENUMBER 912

Hypothesis: POSSIBLE_ALUMINUM

Actions:
CreateObject <POSSIBLE_PEAKS> |POSSIBLE_AL_PEAKS|

As seen above, rules are composed of three parts: conditions, a hypothesis, and actions. Conditions are conditional statements which must be evaluated to a Boolean true or false value. If all of the conditions evaluate to true, the hypothesis, which is also a Boolean value, will evaluate to true, and the actions will be performed. If even one of the conditions evaluates to false, the
hypothesis will evaluate to false, and the actions will not be performed. In the preceding rule, there are two conditions and one action. If both of the conditions evaluate to true, the hypothesis POSSIBLE_ALUMINUM will evaluate to true, and the action will be performed. If the first condition evaluates to false, the hypothesis will automatically evaluate to false and the second will not even be evaluated.

The vertical bars enclosing POSSIBLE_PEAKS indicate that it is a class. The angle brackets outside the vertical bars indicate that pattern matching will occur. The attached property, WAVENUMBER, indicates that the WAVENUMBER slot of each object in the class POSSIBLE_PEAKS will be used for the comparison that is determined by the conditions.

The first condition tells the expert system to pattern match, which means to look at each object (PEAK_1 through PEAK_40) in the class POSSIBLE_PEAKS, and compare the value of the slot PEAK_n.WAVENUMBER to 337. If the value of this slot is greater than 337, the object PEAK_n is kept. If it finds at least one match, the condition will evaluate to true and a list will remain which contains all the objects whose slot PEAK_n.WAVENUMBER is greater than 337. If it doesn't find a match, the condition will evaluate to false, the hypothesis POSSIBLE_ALUMINUM will be set to false and no further conditions will be evaluated. This would indicate that none of the peaks in the class POSSIBLE_PEAKS has a peak location wavenumber that is greater than 337, so they can't possibly be aluminum.
The second condition only gets evaluated if the first condition actually creates a list of objects whose PEAK\_n.WAVENUMBER slot is greater than 337. It says to analyze this list of objects and keep the ones whose PEAK\_n.WAVENUMBER slot is less than 912. If it finds at least one match, the condition will evaluate to true and a list will remain which contains all the objects whose PEAK\_n.WAVE- NUMBER slot is greater than 337 and less than 912. If it doesn't find a match, this condition will evaluate to false which causes the hypothesis POSSIBLE_ALUMINUM to be set to false. This indicates that no peaks were found that fall in the wavenumber range established by the presence of aluminum.

If both conditions evaluate to true, the actions will be performed. A list of peaks will remain which might possibly be aluminum since they fall within its wavenumber range. Therefore, a new class will be created called POSSIBLE_AL_PEAKS. As seen in the example rule, this class will be created with the CreateObject command. The second term after the CreateObject command, |POSSIBLE_AL_PEAKS|, tells the expert system to create the new class. The first term after the CreateObject command, <|POSSIBLE_PEAKS|>, represents the list that was created by the first two conditions. So, the CreateObject command takes the list of objects that was compiled by virtue of the first two conditions and puts it into the new class POSSIBLE_AL_PEAKS.

For example, say the class POSSIBLE_PEAKS contains 100 peaks (PEAK\_1 through PEAK\_100), and the rule PICK_POSSIBLE_ALUMINUM_PEAKS found 15 peaks that fall within
the aluminum wavenumber range. The class POSSIBLE_AL_PEAKS would then contain 15 peaks (Figure 8).

The next step is to analyze the 15 peaks in the class POSSIBLE_AL_PEAKS and determine if the pattern exists which confirms the presence of aluminum. If three of these peaks have wavenumbers corresponding to aluminum lines 1, 2 and 8, then it can be concluded that aluminum is present. The wavenumbers for aluminum lines 1, 2, and 8 are approximately 442.7, 471.9, and 443.9.

A rule must be written that looks at the fifteen slots PEAK_n.WAVENUMBER to determine if there is a wavenumber match with aluminum lines 1, 2, and 8. The name of this rule will be INVESTIGATE_ALUMINUM. It appears on the following page.
The first condition evaluates the status of POSSIBLE_ALUMINUM, which is the hypothesis of the rule 'PICK_POSSIBLE_ALUMINUM_PEAKS.' The operator YES evaluates a Boolean value. This is equivalent to asking "Is the value of the hypothesis POSSIBLE_ALUMINUM true?" If POSSIBLE_ALUMINUM is true, this means that the previous rule found some peaks that might possibly be aluminum and stored them in the class POSSIBLE_AL_PEAKS. These peaks then need to be analyzed to see if the confirming pattern exists. This is accomplished with the rest of the conditions. If the value of the hypothesis POSSIBLE_ALUMINUM is false, no possible aluminum peaks were found, and the class POSSIBLE_AL_PEAKS was never

Name: INVESTIGATE_ALUMINUM

Conditions:

YES POSSIBLE_ALUMINUM

> <|POSSIBLE_AL_PEAKS|>.WAVENUMBER 441.7
< <|POSSIBLE_AL_PEAKS|>.WAVENUMBER 443.7
> <<<POSSIBLE_AL_PEAKS>>>.WAVENUMBER 470.9
< <<<POSSIBLE_AL_PEAKS>>>.WAVENUMBER 472.9
> <<<<<<POSSIBLE_AL_PEAKS>>>>.WAVENUMBER 542.9
< <<<<<<POSSIBLE_AL_PEAKS>>>>.WAVENUMBER 544.9

Hypothesis: ALUMINUM_PRESENT

Actions:

DO <<<POSSIBLE_AL_PEAKS>>>.WAVENUMBER

AL_CALIBRATION_PEAK.WAVENUMBER
created. In this case, the hypothesis ALUMINUM_PRESENT will be automatically set to false, and the rest of the conditions will not be evaluated.

The next six conditions use pattern matching to determine if the aluminum confirmation pattern is present. The first two conditions sort through the list of objects in the class POSSIBLE_AL_PEAKS to see if a peak exists whose PEAK_n.WAVE-NUMBER slot is greater than 441.7 and less than 443.7. Since aluminum line 1 has a peak location of approximately 442.7 wavenumbers, this narrow interval will be sufficient to locate it if present. The next two conditions look for aluminum line 2, whose wavenumber is approximately 471.9. The double set of angle brackets tells the system to pattern match on the original list of peaks in the class POSSIBLE_AL_PEAKS. If only one set of angle brackets was used, this would mean "pattern match on the list created by the first two conditions," which contains only one peak. Since the second two conditions look for a completely different peak, they need to start over again with the original list of objects. The last two conditions employ the same methods to look for aluminum line 3, which has a wavenumber location of 543.9 wavenumbers. Once again, the triple set of angle brackets means to pattern match on the original list, not the list created by the second set of conditions with the double set of angle brackets.

If all the conditions evaluate to true, then the presence of aluminum is confirmed and the hypothesis ALUMINUM_PRESENT is set to true. If the hypothesis is true, the action will be performed. In this rule, the purpose of the action is to assign the value of the slot
AL_CALIBRATION_PEAK.WAVENUMBER the value contained in the PEAK_n.WAVENUMBER slot that was found with the pattern match that occurred in third and fourth conditions in the rule, which will be approximately 471.9 wavenumbers. This is indicated by the double angle brackets around the class [POSSIBLE_AL_PEAKS]. The 'DO' operator assigns the value of the second argument to the value of the first argument. So, after this action is completed, the slot AL_CALIBRATION_PEAK.WAVENUMBER will contain a value of approximately 471.9 wavenumbers.

This calibration peak is used to calculate the concentration of the impurity. Once the area of the calibration peak is known, it can be multiplied by a calibration factor to determine the concentration. Each impurity has its own calibration peak, and aluminum's happens to be the peak that occurs at approximately 471.9 wavenumbers. So, after the exact value of the AL_CALIBRATION_PEAK.WAVENUMBER slot is obtained, it is passed to the FORTRAN program 'FINDAREA' (see Appendix). This program uses this wavenumber to locate the peak in the raw spectral data that is stored in a common block. Once it locates the peak, it uses the trapezoidal integration method to determine its area. The area is then multiplied by the aluminum calibration factor to determine the aluminum concentration in the silicon sample.

If the aluminum confirmation pattern has been identified, the next step is to identify all other lines in the spectrum that are caused by the presence of aluminum. Aluminum may cause as many as 20 different peaks to appear, so 20 different rules are needed for their confirmation. Since the wavenumbers where these peaks occur
is already known, pattern matching will be used on the class |
POSSIBLE_ALUMINUM_PEAKS| to see if they exist. This same

technique that was used to identify the aluminum confirmation

pattern will now be used to identify the individual lines.

The first rule will be called ALUMINUM_LINE_1, and it will

attempt to confirm the presence of the first aluminum line, which

has a peak location of approximately 442.7 wavenumbers. The first

thing this rule must do is check to see if the presence of aluminum

has been confirmed. This is done by examining the value of the

hypothesis ALUMINUM_PRESENT, which the previous rule set to true

if the aluminum confirmation pattern was found. If ALUMINUM_-

PRESENT is true, then the rule must examine all peaks in the class

|POSSIBLE_ALUMINUM_PEAKS| to see if one has a peak location of 442.7

wavenumbers. If so, the slot AL_LINE_1.EXIST will be set to true

and the slot AL_LINE_1.WN will be set to 442.7.

Name: AluminumLine_1

Conditions:

Yes ALUMINUM_PRESENT

> |POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER 442

< |POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER 443.5

Hypothesis: ALUMINUM_LINE

Actions:

Let AL_LINE_1.EXIST TRUE

Do |POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER AL_LINE_1.WN
Twenty rules like this will exist for each possible aluminum line that may be present. For example, aluminum_line_X3 has a peak location of approximately 404 wavenumber. The rule to find this peak is:

Name: Aluminum_Line_X3

Conditions:
Yes ALUMINUM_PRESENT
> <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER 403.5
< <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER 404.5

Hypothesis: ALUMINUM_LINE

Actions:
Let AL_LINE_X3.EXIST TRUE
Do <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER AL_LINE_X3.WN
3.3 PROGRAM FLOW

The previous chapters demonstrated how rules are structured and how they interact. Now it can be shown how these rules fit into the flow of the total program.

From now on, the rules will be shown in the way they appear in the rule network, which is a graphic display of the interrelation of all the rules. Looking at this 'rule tree' allows the developer to analyze the flow of the program. The previous rules were shown in the context of the rule editor, which is where they are created.

For example, the first rule the expert system evaluates is named START_THINGS_OFF:

Yes PEAKS_LOADED
Yes PEAKS_EXAMINED START_THINGS_OFF ANALYZED
Yes PEAKS_ANALYZED

This rule has three conditions on the left-hand side that must evaluate to TRUE before the hypothesis ANALYZED can be set to TRUE. These three conditions represent three separate branches of the rule network. The purpose of this rule is to force the evaluation of these three different branches.

The first branch is represented by the hypothesis PEAKS_LOADED (Figure 9). This branch of the network loads the spectrum and the peak table. After the spectrum and peak table have been loaded, the second branch of the network, represented by the
hypothesis PEAKS_EXAMINED, is put into action (Figure 10). This branch of the network looks at the peak table and classifies the peaks into what they might possibly be, i.e., aluminum, boron, gallium, indium, carbon, oxygen, or unknown. The next branch of the network is represented by the hypothesis PEAKS_ANALYZED (Figure 11). This branch takes the categories of possible peaks that were just created and looks for the confirming patterns that tell which impurities are present. The following pages illustrate how these three branches are connected to the rule START_THINGS_OFF.

The three branches of the rule network interact in the following manner. The three conditions on the left hand side point to the rule name, START_THINGS_OFF, and the rule name points to the hypothesis ANALYZED. To get the process going, the hypothesis ANALYZED must be suggested. Suggesting the hypothesis forces the expert system to determine if its Boolean value is TRUE or FALSE. To do this, it must evaluate the conditions on the left-hand side. ANALYZED will be set to true if all three conditions evaluate to true. If any of the conditions evaluate to false, ANALYZED will be set to false.

In this rule, the left-hand side conditions involve three hypotheses to other rules. PEAKS_LOADED, PEAKS_EXAMINED, and PEAKS_ANALYZED are all hypotheses, and the Yes operator forces the evaluation of their Boolean value. When the expert system attempts to evaluate a hypothesis that appears in a condition, an event called backward chaining takes place which causes other rules in that branch of the rule network to be evaluated. If the value of
Figure 9. The rule START_THINGS_OFF and the branch of the rule network triggered by the hypothesis PEAKS_LOADED.
Figure 10. The PEAKS_EXAMINED branch of the rule network.
Figure 11. The PEAKS_ANALYZED branch of the rule network.
the hypothesis is unknown, then the rule leading to that hypothesis must be evaluated.

Since all the rules in a branch are connected and dependent on each other, trying to evaluate one rule will cause all rules leading up to it to also be evaluated. For example, the first condition in the rule START_THINGS_OFF attempts to determine the value of the hypothesis PEAKS_LOADED, which doesn't yet have a value. To determine its value, the expert system must evaluate the rule LOAD_PEAK_TABLE, since this rule has PEAKS_LOADED as its hypothesis. If all the conditions on the left-hand side of the rule LOAD_PEAK_TABLE evaluate to true, then the hypothesis PEAKS_LOADED will be set to true.

However, the first condition of the rule LOAD_PEAK_TABLE attempts to evaluate the value of SPECTRA_ID_ASKED which is the hypothesis of the rule GET_SPECTRA_ID. So the rule GET_SPECTRA_ID must be evaluated before the hypothesis SPECTRA_ID_ASKED will have a value. Since none of the conditions in the rule GET_SPECTRA_ID attempt to evaluate the hypothesis of another rule, this is the end of the 'backward chain.'

To summarize, 'suggesting' the hypothesis ANALYZED forced the evaluation of the rule START_THINGS_OFF. The first condition in this rule, 'Yes PEAKS_LOADED', forced the evaluation of the rule LOAD_PEAK_TABLE. The first condition of this rule, 'Yes SPECTRA_ID_ASKED', forced the evaluation of the rule GET_SPECTRA_ID. Backward chaining forced the evaluation of all the rules in the PEAKS_LOADED branch of the rule network.
After backward chaining forces the evaluation of the three branches of the rule network, ANALYZED will then be set to true. At this point, the peak table has been loaded. It has been examined so that the peaks are classified as to what impurity they might possibly represent, and the groups of possible peaks have been analyzed to see if the confirmation pattern for the various impurities exist.

If it has been determined that the confirmation pattern exists for one or more impurities, the next step is to identify all the peaks that exist due to their presence. This is accomplished with six more rules, which are named AL_LINES, BO_LINES, GA_LINES, IN_LINES, CA_LINES, and OX_LINES. For example, if aluminum is present, then the rule AL_LINES will be evaluated to identify all of the aluminum peaks. This rule will not be evaluated if aluminum is not present. There are approximately 20 different peaks that can appear for each impurity, and these rules will identify any of these peaks that may be present.

These rules are evaluated with a process called forward chaining, which is the opposite of backward chaining. In backward chaining, when a hypothesis appears as data in a left-hand side condition of some initial rule, the expert system goes backward through the rule network to evaluate all rules that are needed to determine the value of this hypothesis. In forward chaining, once the value of a hypothesis or any other data value is determined, this value is volunteered forward to any other rules that uses it. Any time the expert system gives a rule a data value it needs to evaluate, the rule will automatically get evaluated without having to suggest its hypothesis.
Figure 12 illustrates the concept of forward chaining. The previous example showed how three different branches of the rule network branched backwards from the rule START_THINGS_OFF. This example shows how another section of the rule network branches forward from the same rule. The six rules that identify the peaks of the six impurities depend on the value of the hypothesis ANALYZED of the rule START_THINGS_OFF. This illustration shows how these six rules branch forward from the hypothesis ANALYZED.

After the rule START_THINGS_OFF has been successfully evaluated, the value for ANALYZED will be TRUE. This value will then be volunteered to the six rules AL_LINES, BO_LINES, GA_LINES, IN_LINES, CA_LINES, and OX_LINES. If, for example, the presence of aluminum has been confirmed, the value of the hypothesis ALUMINUM_PRESENT will be TRUE. ALUMINUM_PRESENT is the hypothesis of the rule INVESTIGATE_ALUMINUM, which looked for presence of the aluminum confirmation pattern. So, looking at the rule AL_LINES, the first condition 'Yes ANALYZED' will be true, the second condition 'Yes ALUMINUM_PRESENT' will be true, therefore the hypothesis FIND_AL_LINES will evaluate to TRUE, so the action statement 'Do ALUMINUM_LINE ALUMINUM_LINE' will be performed.

ALUMINUM_LINE is a hypothesis, so the action statement 'Do ALUMINUM_LINE ALUMINUM_LINE' is the same as 'suggesting' this hypothesis. This simply forces the evaluation of any rule that has ALUMINUM_LINE as its hypothesis. In Figure 13 there are twenty rules that have ALUMINUM_LINE as its hypothesis.
Figure 12. Forward chaining from the hypothesis ANALYZED.
Each one of these rules tries to identify a particular peak and sets the value of the slot AL_LINE_n.EXIST to TRUE if that peak is found. It also sets the value of AL_LINE_n.WN to the wavenumber of the peak location. So, the action statement 'Do ALUMINUM_LINE ALUMINUM_LINE' triggers the evaluation of 20 rules which will positively identify all of the peaks that are in the spectrum due to the presence of aluminum.

Following the rules that identify the aluminum lines are the rules that identify the lines for all the other impurities (Figures 14 through 17). It is not possible to show all of the rules for each of the impurities. For the rules that are not shown in their entirety, their peak locations can be found in Table 2.
Figure 13. The aluminum rules.
Figure 14. The boron rules.
Figure 15. The gallium rules.
Figure 16. The indium rules.
Figure 17. The carbon and oxygen rules.
4.0 SAMPLE RUN

Table 3 is an example of the report that is produced by the expert system. For this sample run, it was given a silicon spectrum that contained the impurities boron, gallium, and oxygen. As seen in the report, the expert system was able to identify these impurities, determine their concentrations, and list their lines that were present.
Table 3. The report generated by the expert system.

RESULTS OF IMPURITY CLASSIFICATION
11/13/1990

SPECTRUM NAME: siga819ca
TYPE: abs

------------------------------------------

IMPURITY: BORON
CONCENTRATION: 2.72301772882080081e15

PEAKS DETECTED

<table>
<thead>
<tr>
<th>LINE</th>
<th>WAVENUMBER</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>244.96</td>
</tr>
<tr>
<td>2</td>
<td>278.23</td>
</tr>
<tr>
<td>2p</td>
<td>669.0</td>
</tr>
<tr>
<td>3</td>
<td>309.5</td>
</tr>
<tr>
<td>4</td>
<td>320.08</td>
</tr>
<tr>
<td>8</td>
<td>346.33</td>
</tr>
</tbody>
</table>

------------------------------------------

IMPURITY: GALLIUM
CONCENTRATION: 3.8677543243408203e16

PEAKS DETECTED

<table>
<thead>
<tr>
<th>LINE</th>
<th>WAVENUMBER</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>469.85</td>
</tr>
<tr>
<td>2p</td>
<td>897.04</td>
</tr>
<tr>
<td>3</td>
<td>541.96</td>
</tr>
<tr>
<td>3p</td>
<td>921.13</td>
</tr>
<tr>
<td>4</td>
<td>550.03</td>
</tr>
<tr>
<td>6</td>
<td>569.19</td>
</tr>
<tr>
<td>8</td>
<td>574.21</td>
</tr>
<tr>
<td>10</td>
<td>582.28</td>
</tr>
<tr>
<td>X1</td>
<td>346.33</td>
</tr>
</tbody>
</table>

------------------------------------------

IMPURITY: OXYGEN
CONCENTRATION: 2.581750259399414050e16

PEAKS DETECTED

<table>
<thead>
<tr>
<th>LINE</th>
<th>WAVENUMBER</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1136.92</td>
</tr>
</tbody>
</table>
5.0 TESTING THE EXPERT SYSTEM

In deciding what should be considered in testing the expert system, four main criteria were settled upon:

1. **Easy to use:** The system should be easy to use with a minimum of experience needed in computer use.

2. **Requires little analysis experience:** The expert system should produce satisfactory results when being used by a lower level researcher such as a technician, co-op student or entry-level researcher.

3. **Produces accurate results:** The results generated by the expert system should agree with those of an experienced spectroscopist. These results are often quite subjective and vary considerably when the analysis is performed by different researchers. The expert system should find all the impurities present in a given sample spectra and the calculated impurity concentrations should be within the experimental tolerances that are considered acceptable in this type of spectral analysis.

4. **Produces consistent results:** The expert system should give consistent results if given identical spectra. This may seem like an obvious requirement, but the user is allowed some interaction with the system and can add peaks to the peak table if he feels they are significant when viewing the spectrum.
5.1 DISCUSSION

The expert system was first tested by an experienced spectroscopist with little computer experience. No significant difficulties were experienced by this user. After one tutorial session, the researcher was confident enough to run the system without assistance. A co-op student was the next test subject and required only one short tutorial session to become confident enough to use the system without assistance. The only interaction required by the expert system is for the user to supply a spectrum name and indicate whether it is an absorption coefficient or photoconductivity spectrum. Then the user is asked to examine the spectrum on the screen. This spectrum has all peaks marked that the expert system finds. The user may mark additional peaks if he feels they are significant. If these peaks are simply noise or some other irregularity in the spectrum, then the expert system ignores them. This is all the user is required to input to the system.

In order to test the accuracy of the expert system, its results were compared with the results of a study conducted several years earlier. This study examined the relationship between the area under a specific gallium peak in the spectrum and the gallium concentration for that sample as determined by other means such as Hall Analysis or Photoluminesence. The researcher conducting this study ran a series of gallium doped samples and carefully measured what he felt the gallium dopant-concentration peak areas were. The results of this study were used to calculate a concentration
coefficient relating a specific gallium peak's area to the gallium concentration present in the sample.

Fourteen spectra were analyzed by the expert system in this test. Some runs were made by us, the designers, others by a co-op student working in the lab. The results are displayed in table 4.

Table 4. Results of gallium concentration study test.

<table>
<thead>
<tr>
<th>SAMPLE ID</th>
<th>GALLIUM CONCENTRATION ERROR ( % )</th>
</tr>
</thead>
<tbody>
<tr>
<td>GA0102AC</td>
<td>-4.33</td>
</tr>
<tr>
<td>GA044BAC</td>
<td>-5.66</td>
</tr>
<tr>
<td>GA1004AC</td>
<td>-3.23</td>
</tr>
<tr>
<td>GA1069BAC</td>
<td>-1.46</td>
</tr>
<tr>
<td>GA1305AAC</td>
<td>-7.0</td>
</tr>
<tr>
<td>GA203AC</td>
<td>+0.6</td>
</tr>
<tr>
<td>GA370AC</td>
<td>+2.1</td>
</tr>
<tr>
<td>GA402AC</td>
<td>-6.6</td>
</tr>
<tr>
<td>GA757DAC</td>
<td>+0.1</td>
</tr>
<tr>
<td>GA757EAC</td>
<td>-2.5</td>
</tr>
<tr>
<td>GA757FAC</td>
<td>+0.42</td>
</tr>
<tr>
<td>GA987AC</td>
<td>+1.25</td>
</tr>
<tr>
<td>GA987BAC</td>
<td>-8.96</td>
</tr>
<tr>
<td>GA1072AC</td>
<td>-5.26</td>
</tr>
</tbody>
</table>

The expert system's results for impurity or dopant concentration averaged 2.9% less than the values calculated by the spectroscopist in this study. This figure is well within a reasonable value if comparing the results of two experienced researchers. Typically an error of 20% is not unreasonable due to differences that
occur when determining the area of the calibration peak. A calibration peak is a peak occurring in a spectrum that is always generated by the presence of a particular dopant or impurity. The area of this peak is multiplied by a scale factor to calculate that particular dopant's concentration. If the individual chooses different peak boundaries, then different impurity or dopant concentrations will result. In a noisy spectrum, peak boundaries can be difficult to discern. This situation also occurs when the concentrations are very low and the peak is relatively weak. This can be seen in Figure 18; a typical weak peak occurring in a noisy area of the spectrum. Defining the limits of a peak in a noisy area can be difficult at best. This is where the differences generally occur when two researchers analyze the same spectrum.

The dopant or impurity peaks were classified correctly in all cases. The expert system always detected the presence of all impurities known to be present in the samples. In summary, there was good agreement between the system’s analysis and those of other experienced researchers.
Figure 18. Possible peak boundaries in a noisy spectrum.
6.0 FUTURE PLANS FOR THE EXPERT SYSTEM

The expert-system is completed for use with high purity silicon materials. As stated in previous sections, the scope of the project was limited to silicon based electronic materials. The body of knowledge for these materials was well established. To have included other types of optical materials in the knowledge base would have required more man-hours of development and testing than could be justified. The main goal was to use this as a learning experience for future expert-system work within our laboratory. The completed system is dependable and could be used on a day-to-day basis for research as well as quality-control applications. Unfortunately, silicon based electronic materials are no longer the current materials of interest, so it is unlikely the system will be heavily used.

The next phase of this project is to design a knowledge base for analyzing superlattice materials grown by Molecular Beam Epitaxy (MBE). There currently is not a body of knowledge to draw upon for constructing this new system. When a sufficient number of samples of this type have been analyzed, a set of rules should emerge which will be the basis for the next phase. There has been discussion of designing a closed-loop system where the expert system can control the spectrometer as well as the sample chamber temperature, but this plan is still tentative and requires much more study as well as the full support of the researchers in this area.
The expert-system calls several FORTRAN subroutines to perform routine analysis tasks. These routines are described below.

**FILLSPECTRA** The spectral array of data points is stored in a named Fortran common block. FILLSPECTRA is passed the name of the spectra to be analyzed. It simply locates the spectra data file, opens it and loads the common block with the data points from this file. These data points are subsequently used for the interactive graphics in PEAKPICK and in calculations and graphics in FINDAREA.

**PEAKPICK** This is an interactive graphics routine which plots the subject spectrum on the workstation screen. All peaks that have been located by the FTIR data station are marked and the user is then asked if he wishes to mark additional peaks of interest. By using the attached mouse and cursor all peaks of interest that were not marked by the program can be marked by the user interactively. When complete this routine transfers to a subroutine called REFORM which creates an accompanying table of peaks from the spectrum under scrutiny. This table is in a format readable by the expert-system program. When the knowledge base is loaded this peak table is used as a basis for generating preliminary conclusions about the sample.
FINDAREA The impurity concentration is calculated by multiplying the area of a certain peak associated with that impurity by a calibration factor. The resulting impurity concentration is in the units of atoms/cm\(^2\). Since these peaks are superimposed on a background signal that is often noisy, or the peak itself is very weak, defining the peak limits can be a subjective process. Different researchers can analyze the same spectra and obtain peak areas differing by 20%. This routine, after having been passed the wavenumber of an impurity peak of interest, displays suggested peak boundaries and asks the user if he agrees: if yes then calculations continue based on these peak limits. If no, the user indicates with the mouse and cursor what the peak boundaries should be and these are used for subsequent calculations. The peak area is then calculated by a trapezoidal integration method. The user interaction is handled by a subroutine called PEAKPLOT, called from FINDAREA. The calculated value of the area of the peak of interest is then passed back to the expert-system to be recorded and used in impurity concentration calculations. All FORTRAN callable screen graphics are produced by a graphics software package called CA-DISSPLA, distributed by Computer Associates Inc.
This routine reads in the spectrum points and loads them into a common block called BLOCK1.

INTEGER FUNCTION FILLSPECTRA(THESTR,NATOMS,THEATOMS)
INCLUDE 'NXP$LIBRARY:NXPVMSDEF.FOR'
CHARACTER(*) THESTR
INTEGER*4 THEATOMS(*)
INTEGER*4 RETVAL
LOGICAL FLAG I
if returned TRUE then no errors
INTEGER*4 NATOMS
CHARACTER ANS
CHARACTER*80 LINE
CHARACTER*10 INFILE !The datafile with no extension
CHARACTER*15 DATAFILE ! the spectra file
COMMON /BLOCK1/X(20000), Y(20000), NUMBPTS, INFILE, FLAG ! the spectral array of X-Y pairs

DATAFILE = INFILE//.MAS'
OPEN (FILE=DATAFILE, STATUS='OLD', UNIT=1, CARRIAGECONTROL='LIST', 1 ERR=5000)

the following section reads the input file's header section if there is one
READ(1,30) LINE
30 FORMAT(A80)
DO WHILE (LINE(1:2).EQ.'##')
   READ(1,30) LINE
END DO

get the actual data X-Y pairs
I=1
80       READ(1,*,END=95) X(I), Y(I)
        I=I+1
        NUMBPTS = I
        GO TO 80      ! read until an EOF is found
95       CLOSE( UNIT=1 )    ! normal exit point
        FILLSPECTRA = 1    ! return success
        RETURN
5000     print  5001
5001     format(' BAD FILE NAME ')  return
       END
INTEGER FUNCTION PEAKPICK()

This routine allows the user to view the spectrum with the peaks marked. The user can then interactively mark additional peaks that he feels should be included in the analysis of the sample.

Variable Declaration

**\begin{tabular}{ll}
X,Y & The x & y values of the whole spectra \\
Numbpts & The number of points in the spectra \\
Num & 1/4 of the number of peaks in the whole spectra \\
Npk1-4 & The number of points in each corresponding plot \\
X1,Y1 & The temp var with the points to be plotted \\
for each of the four plots \\
Peak1-4 & The peaks picked by the user for each plot \\
Oldpkx,Oldpky & The peaks found in the .pk file \\
Name & The raw name of the spectra (no exts) \\
Line & Dummy variable used to read in a header line \\
\end{tabular}**

INCLUDE 'NXP$LIBRARY:NXPVMSDEF.FOR'

Common /Block1/X(20000), y(20000), Numbpts, Name,flag

```fortran
Integer Num,Npk1,Npk2,Npk3,Npk4,numold
real X1(5000), Y1(5000)
real Peak1(25,2),Peak2(25,2),Peak3(25,2),Peak4(25,2)
real OldPkx(100), OldPky(100)
character*10 Name
character*15 infile
character*80 Line
logical flag, exist
```

Begin Code

```fortran
IF (EXIST(name//'.pk')) THEN
   Open(unit=1,status='old',file=name//'.pk',err=5000)
   ! Loop to get rid of header in .pk file
   do 20 I=1,6
      read(1,10) line
   10 format(a80)
   20 continue
   ! Loop to read in the old peaks from the .pk file
   I = 1
   30 read(1,* ,end=45) OldPkx(I), OldPky(I)
      I = I + 1
   go to 30
   40 continue
   45 numold = I
```

else

```fortran
```

END
open(unit=1,status='new',file=name//'pk')
do 50 l=1,6
   write(1,*), 'Header Line'
50 continue
Numold = 0
endif
C** Here we send the plot to the screen in four different parts, having
C** the user return the peaks he sees on each of the four plots

Num = Numbpts/4

C****** First Plot
   K=0
   Do 60 l=1,Num
      K=K+1
      X1(k)=x(l)
      Y1(k)=Y(l)
   60 continue
   call Plot(X1,Y1,K,Name,Peak1,Npk1,oldpkx,oldpky,numold)

C****** Second Plot
   K=0
   Do 70 l=(Num-100),(Num*2)
      K=K+1
      X1(k)=x(l)
      Y1(k)=Y(l)
   70 continue
   call Plot(X1,Y1,K,Name,Peak2,Npk2,oldpkx,oldpky,numold)

C****** Third Plot
   K=0
   Do 80 l=(Num*2-100),(Num*3)
      K=K+1
      X1(k)=x(l)
      Y1(k)=Y(l)
   80 continue
   call Plot(X1,Y1,K,Name,Peak3,Npk3,oldpkx,oldpky,numold)

C****** Fourth Plot
   K=0
   Do 90 l=(Num*3-100),Numbpts
      K=K+1
      X1(k)=x(l)
      Y1(k)=Y(l)
   90 continue
   call Plot(X1,Y1,K,Name,Peak4,Npk4,oldpkx,oldpky,numold)
call Donepl !finished plotting
CALL SETDEV(IERR,0)
close(1) !close the .pk file
C**** Now we take the four peak arrays and append them to the Name.pk
C**** file that already exist

infile = name/.pk'
IF (EXIST(infile)) then
    Open(unit=10,status='old', access='append', file=infile)
ELSE
    Open(unit=10,status='new', file=infile)
END IF

do 500 l=1,Npk1
   Write(10,*) peak1(l,1), peak1(l,2)
500 Continue

do 510 l=1,Npk2
   Write(10,*) peak2(l,1), peak2(l,2)
510 Continue

do 520 l=1,Npk3
   Write(10,*) peak3(l,1), peak3(l,2)
520 Continue

do 530 l=1,Npk4
   Write(10,*) peak4(l,1), peak4(l,2)
530 Continue

close(10) !close the .pk file
PEAKPICK = 1 !return success code
CALL REFORM !make a new .nxp peak table
return

c 5000 write(*,*) name/.pk', 'error opening that file!!'
FLAG = .FALSE.
call exit
derib

C *************** PLOT ***********************
subroutine Plot(X,Y,N,Name,Peak,NumPts,oldpKx,oldpky,numold)

real X(5000), Y(5000), Peak(100), npkx(100), npky(100)
real Xmin,Xstep,Xmax,Ymin,Ystep,Ymax
Integer N, NumPts, NumOld
Character*10 Name

N=N-1
call PGPX
   call newclr('whit')
call hwspec(0.0, 'BACK')
call page(13,11.)
call area2d(12,10)
call NEWCLR('viol')

63
call Header(name,100,1.5,1)
call Rndlin(X,N,11.5,Xmin,Xstep,Xmax)
call Rndlin(Y,N,9.5,Ymin,Ystep,Ymax)
call Xname('Wave Number',100)
call Yname('Range',100)
call Graf(Xmin,Xstep,Xmax,Ymin,Ystep,Ymax)
call Newclr('red')
call Curve(X,Y,N,0)
call Newclr('Green')
call Marker(13)
call Setdev(0,0)
call Curve(oldpkx,oldpky,Numold,-1)
Call msgag('Select peaks with left mouse button.',37.1.0,9.75)
Call Message('Click middle button to continue.',31.7.5,9.75)
call Read_Screen(Xmin,Xmax,Ymin,Ymax,Peak,Numpts)
call Endpl(0)
return
end

C************************************************************************** Subroutine Exist
**************************************************************************

Function Exist(FileName)
C
C Tests for the existance of a filename.
C
Character*15 FileName
Integer IERR
Logical Exist
C
C Test for file at CurDate
C
OPEN (File = FileName,status = 'OLD',Unit = 1,iostat = ierr)
If (IERR.EQ.29) then
   Exist = .FALSE.
else
   Exist = .TRUE.
end if
C
Close(Unit = 1)
Return
End

C************************************************************************ Subroutine Read_screen
************************************************************************

subroutine read_screen (xmin,xmax,ymin,ymax,Peak,NumPts)
real xposIx position read from the screen
real yposIy position read from the screen
integer iInumber of points read form the screen
real save_xpos(100)Ithe numbers inputed from the screen

64
real save_ypos(100) ! the numbers inputed from the screen
logical first ! is this the first pos read from the screen?
integer rkey ! status of the return key (0=ok, 1=stop)
real Peak(25,2)
Integer NumPts

C Use the mouse to get the positions from the screen
C
call angle(9.0)
first = .true.
rkey = 0
I = 0
do while (rkey.ne. 1)
  continue
  call get-position(xmin,xmax,ymin,ymax,first,
      &
      xpos,ypos,rkey)
  if (rkey.eq. 0) then
    call MARKER(13)
call CURVE(Xpos,Ypos,1,-1)
    I = I + 1
    Peak(I,1) = Xpos
    Peak(I,2) = Ypos
    save_xpos(I) = xpos
    save_ypos(I) = ypos
go to 10
  else if (rkey.eq. 1) then
    NumPts = I
    return
  else
    type 300
      format(' No valid data available')
  endif
endo
dcall ANGLE(0.0)
return
end

C******************               get_position               ******************

subroutine GET_POSITION(xmin,xmax,ymin,ymax,
&
   first,xpos,ypos,rkey)
Integer rkey ! status return key (0=ok, 1=stop)
Real xmin,xmax,ymin,ymax
Real pos,Ypos ! returned values from the plot
Logical First
Integer IKEY_R
Real Initx, Inity ! initial cross-hair position

if (first) then
call CRDSYS('USER')
   initx=(xmax-xmin)/2.0

\[ \text{infty} = (\text{ymax} - \text{ymin})/2.0 \]
\[ \text{firsts} = \text{.false.} \]
endif
call INILOC(1, xpos, ypos, 0, 0)
call ECHLOC(1, 2, xmin, xmax, ymin, ymax)
call REQLOC(1, rkey, xpos, ypos, ikey_r)

return
end
FINDAREA

This routine is passed the left and right x-coordinates of a peak in a spectral data file. This subroutine calculates the area of that peak and returns it to the expert system for later use.

SUBROUTINE FINDAREA(THESTR,NATOMS,THEATOMS)
INCLUDE 'NXP$LIBRARY:NXPVMSDEF.FOR'
CHARACTER(*) THESTR
INTEGER*4 THEATOMS(*),STAT
INTEGER*4 RETVAL,RETVAL1,RETVAL2,RETVAL3,IDFLAG
LOGICAL FLAG  ! if returned TRUE then no errors
INTEGER*4 NATOMS

COMMON /BLOCK1/X(20000), Y(20000),LOADED, NUMBPTS
! the spectral array of X-Y pairs
CHARACTER*80 LINE
CHARACTER*15 INFILE  ! the spectra file
CHARACTER*15 INCODE
CHARACTER*2 CODE
REAL LASTX,MR,ML,L_EDGE,PEAK
INTEGER DELTA,RINDEX

RETVAL1=NXP$GETATOMINFO(THEATOMS(1),NXPS$_AINFO_VALUE,0,0,
$ ',INFILE,NUMB1 ) ! the spectra file name

RETVAL2=NXP$GETATOMINFO(THEATOMS(2),NXPS$_AINFO_VALUE,0,0,
$ NXPS$_DESC_FLOAT,PEAK,NUMB2 ) ! the peak location

RETVAL3=NXP$GETATOMINFO(THEATOMS(3),NXPS$_AINFO_VALUE,0,0,
$ ',INCODE,NUMB3 ) ! the type of dopant code

IF (RETVAL1 .NE. NXPS$_ERR_NOERR) GO TO 5000
IF (RETVAL2 .NE. NXPS$_ERR_NOERR) GO TO 5000
IF (RETVAL3 .NE. NXPS$_ERR_NOERR) GO TO 5000

CODE= INCODE(1:NUMB3) ! cut off excess stuff

IYPEAKX=IFIND(PEAK) ! get set up for loop
IRIGHT= 0
LEFT= 10
MR = -20
ML= 20

This loop works its way down both sides of the peak, computing the
average slope on either side. When one side bottoms out (slope is
zero) it stops.

```
DO WHILE((ML.GT.0) .AND. (MR.LT.0))
MR= SLOPE(IPEAKX+IRIGHT)  ! get the right side slope
ML= SLOPE(IPEAKX-LEFT)    ! get the left side slope
LEFT= LEFT+1               ! farther down left
IRIGHT= IRIGHT+1           ! farther down right
END DO
```

```
IF (ML.LE.0) DELTA= LEFT + 6     ! why 6 and 4, you'll have to
IF (MR.GE.0) DELTA= IRIGHT + 4   ! draw a picture and convince
                                  ! yourself that these work
```

```
LINDEX= IPEAKX - DELTA         ! the left edge index of the peak
RINDEX = IPEAKX + DELTA       ! the right edge index
L_EDGE = X(LINDEX)
R_EDGE = X(RINDEX)
```

```
CALL PEAKPLOT(L_EDGE,R_EDGE,CODE) ! let the user look at the peak
                                  ! limits & change them if needed
```

```
CALL PKAREA(L_EDGE,R_EDGE,AREA) ! get the area
```

```
IF (AREA .LE. 0) GO TO 5000      ! if it's a bad area, say so
```

```
IF (CODE.EQ.'AL') THEN           ! if it's an aluminum peak
   RETVAL=NXP$GETATOMID('ALUMINUM_CALIBRATION_PEAK_AREA',
                          1 IDFLAG,NXP$_ATYPE_SLOT) ! get the area address
   IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
```

```
   RETVAL=NXP$GETATOMID('ALUMINUM_CALIBRATION_PEAK_START',
                        1 IDFLAG1,NXP$_ATYPE_SLOT) ! get the I-edge address
   IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
```

```
   RETVAL=NXP$GETATOMID('ALUMINUM_CALIBRATION_PEAK_END',
                        1 IDFLAG2,NXP$_ATYPE_SLOT) ! get the R-edge address
   IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
```

```
volunteer the area value
   RETVAL=NXP$VOLUNTEER(IDFLAG,NXP$_DESC_FLOAT,AREA,
                         1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRATSET))
   IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
```

```
volunteer the I-edge value
   RETVAL=NXP$VOLUNTEER(IDFLAG1,NXP$_DESC_FLOAT,L_EDGE,
                         1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRATSET))
   IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
```

```
volunteer the R-edge value
   RETVAL=NXP$VOLUNTEER(IDFLAG2,NXP$_DESC_FLOAT,R_EDGE,
                         1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRATSET))
   IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
```
1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
    GO TO 95
ENDIF

C IF (CODE.EQ.'CA') THEN ! if it's a carbon peak
    RETVAL=NXP$GETATOMID('CARBON_CALIBRATION_PEAK.AREA',
               1 IDFLAG,NXP$$_ATYPE_SLOT) ! get the area address
    IF (RETVAL .NE. NXP$$_ERR_NOERR) GO TO 5000
C
C    RETVAL=NXP$GETATOMID('CARBON_CALIBRATION_PEAK.START',
               1 IDFLAG1,NXP$$_ATYPE_SLOT) ! get the I-edge address
    IF (RETVAL .NE. NXP$$_ERR_NOERR) GO TO 5000
C
C    RETVAL=NXP$GETATOMID('CARBON_CALIBRATION_PEAK.END',
               1 IDFLAG2,NXP$$_ATYPE_SLOT) ! get the R-edge address
    IF (RETVAL .NE. NXP$$_ERR_NOERR) GO TO 5000
C
C volunteer the area value
    RETVAL=NXP$VOLUNTEER(IDFLAG,NXP$$_DESC_FLOAT,AREA,
               1 JIOR(NXP$$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
    IF (RETVAL .NE. NXP$$_ERR_NOERR) GO TO 5000
C
C volunteer the I-edge value
    RETVAL=NXP$VOLUNTEER(IDFLAG1,NXP$$_DESC_FLOAT,L_EDGE,
               1 JIOR(NXP$$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
    IF (RETVAL .NE. NXP$$_ERR_NOERR) GO TO 5000
C
C volunteer the R-edge value
    RETVAL=NXP$VOLUNTEER(IDFLAG2,NXP$$_DESC_FLOAT,R_EDGE,
               1 JIOR(NXP$$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
    IF (RETVAL .NE. NXP$$_ERR_NOERR) GO TO 5000

C
C IF (CODE.EQ.'O') THEN ! if it's an oxygen peak
    RETVAL=NXP$GETATOMID('OXYGEN_CALIBRATION_PEAK.AREA',
               1 IDFLAG,NXP$$_ATYPE_SLOT) ! get the area address
    IF (RETVAL .NE. NXP$$_ERR_NOERR) GO TO 5000
C
C    RETVAL=NXP$GETATOMID('OXYGEN_CALIBRATION_PEAK.START',
               1 IDFLAG1,NXP$$_ATYPE_SLOT) ! get the I-edge address
    IF (RETVAL .NE. NXP$$_ERR_NOERR) GO TO 5000
C
C    RETVAL=NXP$GETATOMID('OXYGEN_CALIBRATION_PEAK.END',
               1 IDFLAG2,NXP$$_ATYPE_SLOT) ! get the R-edge address
    IF (RETVAL .NE. NXP$$_ERR_NOERR) GO TO 5000
C
C volunteer the area value
    RETVAL=NXP$VOLUNTEER(IDFLAG,NXP$$_DESC_FLOAT,AREA,
1  JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

  c volunteer the l-edge value
  c  RETVAL=NXP$VOLUNTEER(IDFLAG1,NXP$_DESC_FLOAT,L_EDGE,
  c  1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
  c  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

  c volunteer the r-edge value
  c  RETVAL=NXP$VOLUNTEER(IDFLAG2,NXP$_DESC_FLOAT,R_EDGE,
  c  1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
  c  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

  GO TO 95
  ENDIF

C------------------------------------------------------------------------

C IF (CODE.EQ.'BO') THEN ! if it's a boron peak
C get the area address
  c  RETVAL=NXP$GETATOMID('BORON_CALIBRATION_PEAK.AREA',
  c     1 IDFLAG,NXP$_ATYPE SLOT)
  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C  RETVAL=NXP$GETATOMID('BORON_CALIBRATION_PEAK.START',
  C     1 IDFLAG1,NXP$_ATYPE SLOT) ! get the l-edge address
C  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C  RETVAL=NXP$GETATOMID('BORON_CALIBRATION_PEAK.END',
  C     1 IDFLAG2,NXP$_ATYPE SLOT) ! get the R-edge address
C  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C volunteer the area value
  c  RETVAL=NXP$VOLUNTEER(IDFLAG,NXP$_DESC_FLOAT,AREA,
  c     1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C volunteer the l-edge value
  c  RETVAL=NXP$VOLUNTEER(IDFLAG1,NXP$_DESC_FLOAT,L_EDGE,
  C     1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
  C  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C volunteer the r-edge value
  c  RETVAL=NXP$VOLUNTEER(IDFLAG2,NXP$_DESC_FLOAT,R_EDGE,
  c     1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
  c  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

  GO TO 95
  ENDIF

C------------------------------------------------------------------------

C IF (CODE.EQ.'IN') THEN ! if it's an indium peak
C get the area address
  c  RETVAL=NXP$GETATOMID('BORON_CALIBRATION_PEAK.AREA',
  C     1 IDFLAG,NXP$_ATYPE SLOT)
  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C  RETVAL=NXP$GETATOMID('BORON_CALIBRATION_PEAK.START',
  C     1 IDFLAG1,NXP$_ATYPE SLOT) ! get the l-edge address
C  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C  RETVAL=NXP$GETATOMID('BORON_CALIBRATION_PEAK.END',
  C     1 IDFLAG2,NXP$_ATYPE SLOT) ! get the R-edge address
C  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C volunteer the area value
  c  RETVAL=NXP$VOLUNTEER(IDFLAG,NXP$_DESC_FLOAT,AREA,
  c     1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C volunteer the l-edge value
  c  RETVAL=NXP$VOLUNTEER(IDFLAG1,NXP$_DESC_FLOAT,L_EDGE,
  C     1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
  C  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C volunteer the r-edge value
  c  RETVAL=NXP$VOLUNTEER(IDFLAG2,NXP$_DESC_FLOAT,R_EDGE,
  C     1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
  C  IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

  GO TO 95
  ENDIF

C------------------------------------------------------------------------

C IF (CODE.EQ.'IN') THEN ! if it's an indium peak

C get the area address

70
RETVAL=NXP$GETATOMID('INDIUM_CALIBRATION_PEAK.AREA',
 1 IDFLAG,NXP$_ATYPE_SLOT)
IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
C RETVAL=NXP$GETATOMID('INDIUM_CALIBRATION_PEAK.START',
 1 IDFLAG1,NXP$_ATYPE_SLOT)  ! get the I-edge address
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
C RETVAL=NXP$GETATOMID('INDIUM_CALIBRATION_PEAK.END',
 1 IDFLAG2,NXP$_ATYPE_SLOT)  ! get the R-edge address
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
C RETVAL=NXP$VOLUNTEER(IDFLAG,NXP$_DESC_FLOAT,AREA,
 1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$MVSTRAT_SET))
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
c volunteer the I-edge value
C RETVAL=NXP$VOLUNTEER(IDFLAG1,NXP$_DESC_FLOAT,I_EDGE,
 1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$MVSTRAT_SET))
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
c volunteer the R-edge value
C RETVAL=NXP$VOLUNTEER(IDFLAG2,NXP$_DESC_FLOAT,R_EDGE,
 1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$MVSTRAT_SET))
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
C GO TO 95
ENDIF
C IF (CODE.EQ.'GA') THEN  ! if it's a gallium peak
C get the area address
C RETVAL=NXP$GETATOMID('GALLIUM_CALIBRATION_PEAK.AREA',
 1 IDFLAG,NXP$_ATYPE_SLOT)
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
C RETVAL=NXP$GETATOMID('GALLIUM_CALIBRATION_PEAK.START',
 1 IDFLAG1,NXP$_ATYPE_SLOT)  ! get the I-edge address
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
C RETVAL=NXP$GETATOMID('GALLIUM_CALIBRATION_PEAK.END',
 1 IDFLAG2,NXP$_ATYPE_SLOT)  ! get the R-edge address
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
C volunteer the area value
C RETVAL=NXP$VOLUNTEER(IDFLAG,NXP$_DESC_FLOAT,AREA,
 1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$MVSTRAT_SET))
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000
c volunteer the I-edge value
C RETVAL=NXP$VOLUNTEER(IDFLAG1,NXP$_DESC_FLOAT,I_EDGE,
 1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$MVSTRAT_SET))
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

C volunteer the r-edge value
C RETVAL=NXP$VOLUNTEER(IDFLAG2,NXP$_DESC_FLOAT,R_EDGE,
C 1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))
C IF (RETVAL .NE. NXP$_ERR_NOERR) GO TO 5000

GO TO 95
ENDIF

5000 FLAG = .FALSE.  ! only exit here if an error occurred
RETVAL=NXP$GETATOMID('STATUS.FINDAREA',IDFLAG,NXP$_ATYPE_SLOT)
RETVAL=NXP$VOLUNTEER(IDFLAG,NXP$_DESC_INT,FLAG,
1 JIOR(NXP$_VSTRAT_NOFWRD,NXP$M_VSTRAT_SET))

95 CLOSE( UNIT=1 )  ! normal exit point
96 RETURN
END

FUNCTION IFIND(POINT)
C This function returns the x index value of POINT in SPECTRA

COMMON /BLOCK1/X(20000), Y(20000),LOADED,NUMBPTS

I=1
DO WHILE (X(I).LT.POINT)
   I=I+1
END DO
IFIND=I
RETURN
END

FUNCTION SLOPE(J)
C This routine calculates the average slope of a curve segment. It
C currently uses 10 points to average. The parameter J is the index
C number of the leftmost point in the slope averaging 'window'.

COMMON /BLOCK1/X(20000), Y(20000),LOADED,NUMBPTS
REAL M,NUM

M=0.0
DO 10 J=1, 10
   NUM= Y(J+1)- Y(J)
   DENOM= X(J+1)- X(J)
   IF(DENOM.EQ.0) DENOM=OLDENOM
   M= NUM/DENOM + M
   OLDENOM= DENOM
10 CONTINUE
RETURN
END
J = J + 1
CONTINUE
SLOPE = M/10.0
RETURN
END

Subroutine PKAREA(STP, ENP, AREA)

This program, given a set of endpoints and the (x,y) points
in between, will calculate the area under the peak using
a trapezoidal approximation. First calculating the area
from the peak to the zero baseline and then subtracting
from that the area from the zero baseline to the peak baseline.

Variable Declarations

STP -> starting point of the peak
ENP -> ending point of the peak
SPECTRA -> all the points in the spectra represented by
their (x,y) values in the array form:
    (data #, x or y) = array representation
    (1,x) = (1,1)
    (1,y) = (1,2)
    (2,x) = (2,1)
    (2,y) = (2,2)
    : 
    : 
    (n,x) = (n,1)
    (n,y) = (n,2)
RESAR -> area from peak baseline to zero baseline
SUM -> area under the peak and above the zero baseline
AREA -> the area under the peak and above the peak baseline
I -> loop control variables

Common /BLOCK1/X(20000), Y(20000), LOADED, NUMBPTS
Real EP(2), SP(2)

*I Loop to find the starting point in the spectra *

I = 1
Do While (STP .gt. X(I))
    I = I + 1
End Do
I = I - 1
SP(1) = X(I)
SP(2) = Y(I)
**Loop to calculate the area from the peak to the zero baseline**

```
SUM = 0.0

Do While (ENP .ge. X(I))
    SUM = SUM + 0.5 * (X(I+1) - X(I))
    & (Y(I) + Y(I+1))
    I = I + 1
End do

EP(1) = X(I)
EP(2) = Y(I)

**Calculate RESAR and subtract it from SUM**

RESAR = 0.5 * (EP(1) - SP(1))*(EP(2) + SP(2))
AREA = SUM - RESAR
RETURN
END
```
APPENDIX B. LIST OF RULES

RULE: Rule INVESTIGATE_ALUMINUM_ABS (#1)
If
  there is evidence of ALUMINUM_PRESENT
  And SPECTRA.TYPE is "ABS"
Then ALUMINUMANALYZED_ABS
  is confirmed.
  And CODE is set to "AL"
  And Execute
  "FINDAREA"(@ATOMID=SPECTRA.POINTS,ALUMINUM_CALIBRATION_PEAK.WAVENUMBER, CODE.VALUE;)
    And (ALUMINUM_CALIBRATION_PEAK.AREA*4.1666e+13) is assigned to ALUMINUM_CALIBRATION_PEAK.CONC

RULE: Rule INVESTIGATE_ALUMINUM_PHOTO (#2)
If
  there is evidence of ALUMINUM_PRESENT
  And SPECTRA.TYPE is "PHOTO"
Then ALUMINUMANALYZED_PHOTO
  is confirmed.
  And CODE is set to "AL"
  And 0.0 is assigned to ALUMINUM_CALIBRATION_PEAK.AREA

RULE: Rule ALUMINUMLINE_X4 (#3)
If
  there is evidence of ALUMINUM_PRESENT
  And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is greater than 413.5
  And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is less than 414.5
Then ALUMINUM_LINE
  is confirmed.
  And AL_LINE_X4.EXIST is set to TRUE
  And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is assigned to AL_LINE_X4.WN
RULE : Rule ALUMINUM_LINE_X3 (#4)
If
there is evidence of ALUMINUM_PRESENT
And <|[POSSIBLE_ALUMINUM_PEAKS]|>.WAVENUMBER is greater than 403.5
And <|[POSSIBLE_ALUMINUM_PEAKS]|>.WAVENUMBER is less than 404.5
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_X3.EXIST is set to TRUE
And <|[POSSIBLE_ALUMINUM_PEAKS]|>.WAVENUMBER is assigned to AL_LINE_X3.WN

RULE : Rule ALUMINUM_LINE_X2 (#5)
If
there is evidence of ALUMINUM_PRESENT
And <|[POSSIBLE_ALUMINUM_PEAKS]|>.WAVENUMBER is greater than 371.5
And <|[POSSIBLE_ALUMINUM_PEAKS]|>.WAVENUMBER is less than 372.5
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_X2.EXIST is set to TRUE
And <|[POSSIBLE_ALUMINUM_PEAKS]|>.WAVENUMBER is assigned to AL_LINE_X2.WN

RULE : Rule ALUMINUM_LINE_X1 (#6)
If
there is evidence of ALUMINUM_PRESENT
And <|[POSSIBLE_ALUMINUM_PEAKS]|>.WAVENUMBER is greater than 337.5
And <|[POSSIBLE_ALUMINUM_PEAKS]|>.WAVENUMBER is less than 338.5
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_X1.EXIST is set to TRUE
And <|[POSSIBLE_ALUMINUM_PEAKS]|>.WAVENUMBER is assigned to AL_LINE_X1.WN
RULE : Rule ALUMINUM_LINE_EX (#7)
If
there is evidence of ALUMINUM_PRESENT
And <POSSIBLE_ALUMINUM_PEAWS>.WAVENUMBER is greater than 453.5
And <POSSIBLE_ALUMINUM_PEAWS>.WAVENUMBER is less than 4454.5
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_EX.EXIST is set to TRUE
And <POSSIBLE_ALUMINUM_PEAWS>.WAVENUMBER is assigned to AL_LINE_EX.WN

RULE : Rule ALUMINUM_LINE_ESTAR (#8)
If
there is evidence of ALUMINUM_PRESENT
And <POSSIBLE_ALUMINUM_PEAWS>.WAVENUMBER is greater than 911.5
And <POSSIBLE_ALUMINUM_PEAWS>.WAVENUMBER is less than 912
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_ESTAR.EXIST is set to TRUE
And <POSSIBLE_ALUMINUM_PEAWS>.WAVENUMBER is assigned to AL_LINE_ESTAR.WN

RULE : Rule ALUMINUM_LINE_EL (#9)
If
there is evidence of ALUMINUM_PRESENT
And <POSSIBLE_ALUMINUM_PEAWS>.WAVENUMBER is greater than 556.5
And <POSSIBLE_ALUMINUM_PEAWS>.WAVENUMBER is less than 557
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_EL.EXIST is set to TRUE
And <POSSIBLE_ALUMINUM_PEAWS>.WAVENUMBER is assigned to AL_LINE_EL.WN
RULE : Rule ALUMINUM_LINE_9 (#10)
If there is evidence of ALUMINUM_PRESENT
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is greater than 546
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is less than 548
Then ALUMINUM_LINE is confirmed.
And AL_LINE_9.EXIST is set to TRUE
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is assigned to AL_LINE_9.WN

RULE : Rule ALUMINUM_LINE_8 (#11)
If there is evidence of ALUMINUM_PRESENT
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is greater than 543
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is less than 546
Then ALUMINUM_LINE is confirmed.
And AL_LINE_8.EXIST is set to TRUE
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is assigned to AL_LINE_8.WN

RULE : Rule ALUMINUM_LINE_7 (#12)
If there is evidence of ALUMINUM_PRESENT
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is greater than 540.7
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is less than 542
Then ALUMINUM_LINE is confirmed.
And AL_LINE_7.EXIST is set to TRUE
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is assigned to AL_LINE_7.WN
RULE : Rule ALUMINUM_LINE_6 (#13)
If
there is evidence of ALUMINUM_PRESENT
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is greater than 538
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is less than 539.3
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_6.EXIST is set to TRUE
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is assigned to AL_LINE_6.WN

RULE : Rule ALUMINUM_LINE_5 (#14)
If
there is evidence of ALUMINUM_PRESENT
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is greater than 534.5
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is less than 535.5
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_5.EXIST is set to TRUE
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is assigned to AL_LINE_5.WN

RULE : Rule ALUMINUM_LINE_4P (#15)
If
there is evidence of ALUMINUM_PRESENT
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is greater than 900
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is less than 900.5
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_4P.EXIST is set to TRUE
And <POSSIBLE_ALUMINUM_PEAKS>.WAVENUMBER is assigned to AL_LINE_4P.WN

79
RULE : Rule ALUMINUM_LINE_3P (#16)
If
there is evidence of ALUMINUM_PRESENT
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is greater than 891.5
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is less than 892
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_3P.EXIST is set to TRUE
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is assigned to AL_LINE_3P.WN

RULE : Rule ALUMINUM_LINE_2P (#17)
If
there is evidence of ALUMINUM_PRESENT
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is greater than 867
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is less than 868
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_2P.EXIST is set to TRUE
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is assigned to AL_LINE_2P.WN

RULE : Rule ALUMINUM_LINE_2 (#18)
If
there is evidence of ALUMINUM_PRESENT
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is greater than 471
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is less than 472.5
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_2.EXIST is set to TRUE
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is assigned to AL_LINE_2.WN
RULE : Rule ALUMINUM_LINE_11 (#19)
If
there is evidence of ALUMINUM_PRESENT
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is greater than 554.5
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is less than 556.4
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_11.EXIST is set to TRUE
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is assigned to AL_LINE_11.WN

RULE : Rule ALUMINUM_LINE_10 (#20)
If
there is evidence of ALUMINUM_PRESENT
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is greater than 549
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is less than 554
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_10.EXIST is set to TRUE
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is assigned to AL_LINE_10.WN

RULE : Rule ALUMINUM_LINE_1 (#21)
If
there is evidence of ALUMINUM_PRESENT
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is greater than 442
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is less than 443.5
Then ALUMINUM_LINE
is confirmed.
And AL_LINE_1.EXIST is set to TRUE
And <|POSSIBLE_ALUMINUM_PEAKS|>.WAVENUMBER is assigned to AL_LINE_1.WN
RULE : Rule INVESTIGATE_ALUMINUM (#22)
If
there is evidence of POSSIBLE_ALUMINUM
And <POSSIBLE_ALUMINUM_PEEKS>.WAVENUMBER is greater than 441.7
And <POSSIBLE_ALUMINUM_PEEKS>.WAVENUMBER is less than 443.7
And <<POSSIBLE_ALUMINUM_PEEKS>>.WAVENUMBER is greater than 470.9
And <<POSSIBLE_ALUMINUM_PEEKS>>.WAVENUMBER is less than 472.9
And <<<POSSIBLE_ALUMINUM_PEEKS>>>.WAVENUMBER is greater than 542.9
And <<<POSSIBLE_ALUMINUM_PEEKS>>>.WAVENUMBER is less than 544.9
Then ALUMINUM_PRESENT
is confirmed.
And <<POSSIBLE_ALUMINUM_PEEKS>>.WAVENUMBER is assigned to ALUMINUM_CALIBRATION_PEAK.WAVENUMBER

RULE : Rule START_THINGS_OFF (#23)
If
there is evidence of PEAKS_LOADED
And there is evidence of PEAKS_EXAMINED
And there is evidence of PEAKS_ANALYZED
Then ANALYZED
is confirmed.

RULE : Rule INVESTIGATE_BORON_ABS (#24)
If
there is evidence of BORON_PRESENT
And SPECTRA.TYPE is "ABS"
Then BORON_ANALYZED_ABS
is confirmed.
And CODE is set to "BO"
And Execute
"FINDAREA"(@ATOMID=SPECTRA.POINTS,BORON_CALIBRATION_PEAK.WAVENUMBER,CODE.Value;)
And (BORON_CALIBRATION_PEAK.AREA*1.5151e+13) is assigned to BORON_CALIBRATION_PEAK_CONC

82
RULE : Rule INVESTIGATE_BORON_PHOTO (#25)
If
  there is evidence of BORON_PRESENT
  And SPECTRA.TYPE is "PHOTO"
Then BORON_ANALYZED_PHOTO
  is confirmed.
  And CODE is set to "BO"
  And 0.0 is assigned to BORON_CALIBRATION_PEAK.AREA

RULE : Rule BORON_LINE_X4 (#26)
If
  there is evidence of BORON_PRESENT
  And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is greater than 260.0
  And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is less than 262.0
Then BORON_LINE
  is confirmed.
  And BO_LINE_X4.EXIST is set to TRUE
  And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is assigned to BO_LINE_X4.WN

RULE : Rule BORON_LINE_X3 (#27)
If
  there is evidence of BORON_PRESENT
  And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is greater than 249.0
  And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is less than 251.0
Then BORON_LINE
  is confirmed.
  And BO_LINE_X3.EXIST is set to TRUE
  And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is assigned to BO_LINE_X3.WN
RULE : Rule BORON_LINE_X2 (#28)
If
    there is evidence of BORON_PRESENT
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is greater than 219.0
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is less than 221.0
Then BORON_LINE
    is confirmed.
    And BO_LINE_X2.EXIST is set to TRUE
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is assigned to BO_LINE_X2.WN

RULE : Rule BORON_LINE_X1 (#29)
If
    there is evidence of BORON_PRESENT
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is greater than 183.5
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is less than 184.5
Then BORON_LINE
    is confirmed.
    And BO_LINE_X1.EXIST is set to TRUE
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is assigned to BO_LINE_X1.WN

RULE : Rule BORON_LINE_EX (#30)
If
    there is evidence of BORON_PRESENT
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is greater than 298.0
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is less than 300.0
Then BORON_LINE
    is confirmed.
    And BO_LINE_EX.EXIST is set to TRUE
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is assigned to BO_LINE_EX.WN
RULE : Rule BORON_LINE_ESTAR (#31)
  If
    there is evidence of BORON_PRESENT
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is greater than 712.4
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is less than 713.4
  Then BORON_LINE
    is confirmed.
    And BO_LINE_ESTAR.EXIST is set to TRUE
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is assigned to BO_LINE_ESTAR.WN

RULE : Rule BORON_LINE_E (#32)
  If
    there is evidence of BORON_PRESENT
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is greater than 358.0
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is less than 359.0
  Then BORON_LINE
    is confirmed.
    And BO_LINE_E.EXIST is set to TRUE
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is assigned to BO_LINE_E.WN

RULE : Rule BORON_LINE_9 (#33)
  If
    there is evidence of BORON_PRESENT
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is greater than 348.6
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is less than 349.3
  Then BORON_LINE
    is confirmed.
    And BO_LINE_9.EXIST is set to TRUE
    And <POSSIBLE_BORON_PEAKS>.WAVENUMBER is assigned to BO_LINE_9.WN
RULE: Rule BORON_LINE_8 (#34)
If
   there is evidence of BORON_PRESENT
   And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 344.2
   And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 346.7
Then BORON_LINE
   is confirmed.
   And BO_LINE_8.EXIST is set to TRUE
   And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_8.WN

RULE: Rule BORON_LINE_7 (#35)
If
   there is evidence of BORON_PRESENT
   And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 341.6
   And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 342.6
Then BORON_LINE
   is confirmed.
   And BO_LINE_7.EXIST is set to TRUE
   And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_7.WN

RULE: Rule BORON_LINE_6 (#36)
If
   there is evidence of BORON_PRESENT
   And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 337.5
   And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 340.6
Then BORON_LINE
   is confirmed.
   And BO_LINE_6.EXIST is set to TRUE
   And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_6.WN
RULE : Rule BORON_LINE_5 (#37)
If
there is evidence of BORON_PRESENT
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 334.0
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 335.0
Then BORON_LINE
is confirmed.
And BO_LINE_5.EXIST is set to TRUE
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_5.WN

RULE : Rule BORON_LINE_4P (#38)
If
there is evidence of BORON_PRESENT
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 700.8
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 701.8
Then BORON_LINE
is confirmed.
And BO_LINE_4P.EXIST is set to TRUE
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_4P.WN

RULE : Rule BORON_LINE_4 (#39)
If
there is evidence of BORON_PRESENT
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 318.8
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 322.4
Then BORON_LINE
is confirmed.
And BO_LINE_4.EXIST is set to TRUE
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_4.WN

87
RULE : Rule BORON_LINE_3P (#40)

If

there is evidence of BORON_PRESENT
And <[POSSIBLE_BORON_PEAKS]> . WAVENUMBER is greater than 692.3
And <[POSSIBLE_BORON_PEAKS]> . WAVENUMBER is less than 693.4

Then BORON_LINE
is confirmed.
And BO_LINE_3P.EXIST is set to TRUE
And <[POSSIBLE_BORON_PEAKS]> . WAVENUMBER is assigned to BO_LINE_3P.WN

RULE : Rule BORON_LINE_3 (#41)

If

there is evidence of BORON_PRESENT
And <[POSSIBLE_BORON_PEAKS]> . WAVENUMBER is greater than 308.8
And <[POSSIBLE_BORON_PEAKS]> . WAVENUMBER is less than 309.8

Then BORON_LINE
is confirmed.
And BO_LINE_3.EXIST is set to TRUE
And <[POSSIBLE_BORON_PEAKS]> . WAVENUMBER is assigned to BO_LINE_3.WN

RULE : Rule BORON_LINE_2P (#42)

If

there is evidence of BORON_PRESENT
And <[POSSIBLE_BORON_PEAKS]> . WAVENUMBER is greater than 668.1
And <[POSSIBLE_BORON_PEAKS]> . WAVENUMBER is less than 669.1

Then BORON_LINE
is confirmed.
And BO_LINE_2P.EXIST is set to TRUE
And <[POSSIBLE_BORON_PEAKS]> . WAVENUMBER is assigned to BO_LINE_2P.WN
RULE : Rule BORON_LINE_2 (#43)
If
there is evidence of BORON_PRESENT
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 277.7
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 278.7
Then BORON_LINE
is confirmed.
And BO_LINE_2.EXIST is set to TRUE
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_2.WN

RULE : Rule BORON_LINE_11 (#44)
If
there is evidence of BORON_PRESENT
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 356.0
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 357.6
Then BORON_LINE
is confirmed.
And BO_LINE_11.EXIST is set to TRUE
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_11.WN

RULE : Rule BORON_LINE_10 (#45)
If
there is evidence of BORON_PRESENT
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 350.3
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 356.0
Then BORON_LINE
is confirmed.
And BO_LINE_10.EXIST is set to TRUE
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_10.WN
RULE : Rule BORON_LINE_1 (#46)
If there is evidence of BORON_PRESENT
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 244.5
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 245.5
Then BORON_LINE
is confirmed.
And BO_LINE_1.EXIST is set to TRUE
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is assigned to BO_LINE_1.WN

RULE : Rule INVESTIGATE_BORON (#47)
If there is evidence of POSSIBLE_BORON
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is greater than 244.5
And <|POSSIBLE_BORON_PEAKS|>.WAVENUMBER is less than 245.5
And <|POSSIBLE_BORON_PEAKS|>>.WAVENUMBER is greater than 277.2
And <|POSSIBLE_BORON_PEAKS|>>.WAVENUMBER is less than 279.2
And <|POSSIBLE_BORON_PEAKS|>>>.WAVENUMBER is greater than 318.3
And <|POSSIBLE_BORON_PEAKS|>>>.WAVENUMBER is less than 320.3
Then BORON_PRESENT
is confirmed.
And <|POSSIBLE_BORON_PEAKS|>>.WAVENUMBER is assigned to BORON_CALIBRATION_PEAK.WAVENUMBER
RULE : Rule INVESTIGATE_CARBON_ABS (#48)
If
   there is evidence of CARBON_PRESENT
   And SPECTRA.TYPE is "ABS"
Then CARBON_ANALYZED_ABS
   is confirmed.
   And CODE is set to "CA"
   And Fxecute
   "FINDAREA"(@ATOMID=SPECTRA.POINTS,CARBON_CALIBRATION_PEAK.
   WAVENUMBER,CODE.Value;)
   And (CARBON_CALIBRATION_PEAK.AREA*6.700e+16) is
   assigned to CARBON_CALIBRATION_PEAK.CONC

RULE : Rule INVESTIGATE_CARBON_PHOTO (#49)
If
   there is evidence of CARBON_PRESENT
   And SPECTRA.TYPE is "PHOTO"
Then CARBON_ANALYZED_PHOTO
   is confirmed.
   And CODE is set to "CA"
   And 0.0 is assigned to CARBON_CALIBRATION_PEAK.AREA

RULE : Rule CARBON_LINE_1 (#50)
If
   there is evidence of CARBON_PRESENT
   And <IPossible_Carbon_Peaks>.WAVENUMBER is greater than
   599.5
   And <IPossible_Carbon_Peaks>.WAVENUMBER is less than
   600.5
Then CARBON_LINE
   is confirmed.
   And CA_LINE_1.EXIST is set to TRUE
   And <IPossible_Carbon_Peaks>.WAVENUMBER is assigned to
   CA_LINE_1.WN
RULE: Rule INVESTIGATE_CARBON (#51)
If
    there is evidence of POSSIBLE_CARBON
    And <|POSSIBLE_CARBON_PEAKS|>.WAVENUMBER is greater than 599.5
    And <|POSSIBLE_CARBON_PEAKS|>.WAVENUMBER is less than 600.5
Then CARBON_PRESENT
    is confirmed.
    And <|POSSIBLE_CARBON_PEAKS|>.WAVENUMBER is assigned to CARBON_CALIBRATION_PEAK.WAVENUMBER

RULE: Rule AL_LINES (#52)
If
    there is evidence of ANALYZED
    And there is evidence of ALUMINUM_PRESENT
Then FIND_AL_LINES
    is confirmed.
    And ALUMINUM_LINE is assigned to ALUMINUM_LINE

RULE: Rule BO_LINES (#53)
If
    there is evidence of ANALYZED
    And there is evidence of BORON_PRESENT
Then FIND_BO_LINES
    is confirmed.
    And BORON_LINE is assigned to BORON_LINE

RULE: Rule CA_LINES (#54)
If
    there is evidence of ANALYZED
    And there is evidence of CARBON_PRESENT
Then FIND_CA_LINES
    is confirmed.
    And CARBON_LINE is assigned to CARBON_LINE
RULE : Rule GA_LINES (#55)
If
there is evidence of ANALYZED
And there is evidence of GALLIUM_PRESENT
Then FIND_GA_LINES
is confirmed.
And GALLIUM_LINE is assigned to GALLIUM_LINE

RULE : Rule IN_LINES (#56)
If
there is evidence of ANALYZED
And there is evidence of INDIUM_PRESENT
Then FIND_IN_LINES
is confirmed.
And INDIUM_LINE is assigned to INDIUM_LINE

RULE : Rule OX_LINES (#57)
If
there is evidence of ANALYZED
And there is evidence of OXYGEN_PRESENT
Then FIND_OX_LINES
is confirmed.
And OXYGEN_LINE is assigned to OXYGEN_LINE

RULE : Rule INVESTIGATE_GALLIUM_ABS (#58)
If
there is evidence of GALLIUM_PRESENT
And SPECTRA.TYPE is "ABS"
Then GALLIUM_ANALYZED_ABS
is confirmed.
And CODE is set to "GA"
And Execute
"FINDAREA"(ATOMID=SPECTRA.POINTS,GALLIUM_CALIBRATION_PEAK.WAVENUMBER,CODE.Value;)
And (GALLIUM_CALIBRATION_PEAK.AREA*2.780e+14)
is assigned to GALLIUM_CALIBRATION_PEAK.CONC

93
RULE: Rule INVESTIGATE_GALLIUM_PHOTO (#59)
If
   there is evidence of GALLIUM_PRESENT
   And SPECTRA_TYPE is "PHOTO"
Then GALLIUM_ANALYZED_PHOTO
   is confirmed.
   And CODE is set to "GA"
   And 0.0 is assigned to GALLIUM_CALIBRATION_PEAK_AREA

RULE: Rule GALLIUM_LINE_X4 (#60)
If
   there is evidence of GALLIUM_PRESENT
   And <IPOSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is greater than 422
   And <IPOSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is less than 424
Then GALLIUM_LINE
   is confirmed.
   And GA_LINE_X4.EXIST is set to TRUE
   And <IPOSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is assigned to GA_LINE_X4.WN

RULE: Rule GALLIUM_LINE_X2 (#61)
If
   there is evidence of GALLIUM_PRESENT
   And <IPOSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is greater than 380
   And <IPOSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is less than 382
Then GALLIUM_LINE
   is confirmed.
   And GA_LINE_X2.EXIST is set to TRUE
   And <IPOSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is assigned to GA_LINE_X2.WN
RULE : Rule GALLIUM_LINE_X1 (#62)
If
  there is evidence of GALLIUM_PRESENT
  And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater than 346
  And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than 348
Then GALLIUM_LINE
  is confirmed.
  And GA_LINE_X1.EXIST is set to TRUE
  And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to GA_LINE_X1.WN

RULE : Rule GALLIUM_LINE_EX (#63)
If
  there is evidence of GALLIUM_PRESENT
  And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater than 459
  And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than 461
Then GALLIUM_LINE
  is confirmed.
  And GA_LINE_EX.EXIST is set to TRUE
  And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to GA_LINE_EX.WN

RULE : Rule GALLIUM_LINE_ESTAR (#64)
If
  there is evidence of GALLIUM_PRESENT
  And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater than 941
  And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than 942
Then GALLIUM_LINE
  is confirmed.
  And GA_LINE_ESTAR.EXIST is set to TRUE
  And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to GA_LINE_ESTAR.WN
RULE: Rule GALLIUM_LINE_EI (#65)
If there is evidence of GALLIUM_PRESENT
And [<POSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is greater than 586
And [<POSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is less than 587.2
Then GALLIUM_LINE
is confirmed.
And GA_LINE_EI.EXIST is set to TRUE
And [<POSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is assigned to GA_LINE_EI.WN

RULE: Rule GALLIUM_LINE_9 (#66)
If there is evidence of GALLIUM_PRESENT
And [<POSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is greater than 576
And [<POSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is less than 578.2
Then GALLIUM_LINE
is confirmed.
And GA_LINE_9.EXIST is set to TRUE
And [<POSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is assigned to GA_LINE_9.WN

RULE: Rule GALLIUM_LINE_8 (#67)
If there is evidence of GALLIUM_PRESENT
And [<POSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is greater than 573
And [<POSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is less than 576
Then GALLIUM_LINE
is confirmed.
And GA_LINE_8.EXIST is set to TRUE
And [<POSSIBLE_GALLIUM_PEAKS>.WAVENUMBER is assigned to GA_LINE_8.WN
RULE: Rule GALLIUM_LINE_7 (#68)
If
there is evidence of GALLIUM_PRESENT
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater
than 570.5
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than
571.5
Then GALLIUM_LINE
is confirmed.
And GA_LINE_7.EXIST is set to TRUE
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to
GA_LINE_7.WN

RULE: Rule GALLIUM_LINE_6 (#69)
If
there is evidence of GALLIUM_PRESENT
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater
than 567
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than
569.5
Then GALLIUM_LINE
is confirmed.
And GA_LINE_6.EXIST is set to TRUE
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to
GA_LINE_6.WN

RULE: Rule GALLIUM_LINE_5P (#70)
If
there is evidence of GALLIUM_PRESENT
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater
than 933.5
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than
934.5
Then GALLIUM_LINE
is confirmed.
And GA_LINE_5P.EXIST is set to TRUE
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to
GA_LINE_5P.WN
RULE : Rule GALLIUM_LINE_5 (#71)
If
there is evidence of GALLIUM_PRESENT
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater than 563.7
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than 564.7
Then GALLIUM_LINE
is confirmed.
And GA_LINE_5.EXIST is set to TRUE
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to GA_LINE_5.WN

RULE : Rule GALLIUM_LINE_4P (#72)
If
there is evidence of GALLIUM_PRESENT
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater than 929.5
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than 930.5
Then GALLIUM_LINE
is confirmed.
And GA_LINE_4P.EXIST is set to TRUE
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to GA_LINE_4P.WN

RULE : Rule GALLIUM_LINE_4 (#73)
If
there is evidence of GALLIUM_PRESENT
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater than 547.7
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than 553
Then GALLIUM_LINE
is confirmed.
And GA_LINE_4.EXIST is set to TRUE
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to GA_LINE_4.WN
RULE : Rule GALLIUM_LINE_3P (#74)
If
    there is evidence of GALLIUM_PRESENT
    And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater
    than 921
    And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than
    922
Then GALLIUM_LINE
    is confirmed.
    And GA_LINE_3P.EXIST is set to TRUE
    And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to
    GA_LINE_3P.WN

RULE : Rule GALLIUM_LINE_3 (#75)
If
    there is evidence of GALLIUM_PRESENT
    And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater
    than 541
    And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than
    542
Then GALLIUM_LINE
    is confirmed.
    And GA_LINE_3.EXIST is set to TRUE
    And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to
    GA_LINE_3.WN

RULE : Rule GALLIUM_LINE_2P (#76)
If
    there is evidence of GALLIUM_PRESENT
    And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater
    than 896.7
    And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than
    897.7
Then GALLIUM_LINE
    is confirmed.
    And GA_LINE_2P.EXIST is set to TRUE
    And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to
    GA_LINE_2P.WN
RULE : Rule GALLIUM_LINE_11 (#77)
If
there is evidence of GALLIUM_PRESENT
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater
than 584
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than
586.3
Then GALLIUM_LINE
is confirmed.
And GA_LINE_11.EXIST is set to TRUE
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to
GA_LINE_11.WN

RULE : Rule GALLIUM_LINE_10 (#78)
If
there is evidence of GALLIUM_PRESENT
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater
than 579
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than
584
Then GALLIUM_LINE
is confirmed.
And GA_LINE_10.EXIST is set to TRUE
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to
GA_LINE_10.WN

RULE : Rule GALLIUM_LINE_1 (#79)
If
there is evidence of GALLIUM_PRESENT
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is greater
than 469.25
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is less than
470.25
Then GALLIUM_LINE
is confirmed.
And GA_LINE_1.EXIST is set to TRUE
And <|POSSIBLE_GALLIUM_PEAKS|>.WAVENUMBER is assigned to
GA_LINE_1.WN
RULE: Rule INVESTIGATE_GALLIUM (#80)
If there is evidence of POSSIBLE_GALLIUM
And \{POSSIBLE_GALLIUM_PEAKS\}.WAVENUMBER is greater than 468.8
And \{POSSIBLE_GALLIUM_PEAKS\}.WAVENUMBER is less than 470.8
And \{POSSIBLE_GALLIUM_PEAKS\}.WAVENUMBER is greater than 540.5
And \{POSSIBLE_GALLIUM_PEAKS\}.WAVENUMBER is less than 542.5
And \{POSSIBLE_GALLIUM_PEAKS\}.WAVENUMBER is greater than 547.2
And \{POSSIBLE_GALLIUM_PEAKS\}.WAVENUMBER is less than 549.2
Then GALLIUM_PRESENT is confirmed.
And \{POSSIBLE_GALLIUM_PEAKS\}.WAVENUMBER is assigned to GALLIUM_CALIBRATION_PEAK.WAVENUMBER

RULE: Rule INVESTIGATE_INDIUM_ABS (#81)
If there is evidence of INDIUM_PRESENT
And SPECTRA.TYPE is "ABS"
Then INDIUM_ANALYZED_ABS is confirmed.
And CODE is set to "IN"
And Execute "FINDAREA"(@ATOMID=SPECTRA.POINTS, INDIUM_CALIBRATION_PEAK.WAVENUMBER, CODE.Value;)
And (INDIUM_CALIBRATION_PEAK.AREA*2.0000000000e+15) is assigned to INDIUM_CALIBRATION_PEAK.CONC

RULE: Rule INVESTIGATE_INDIUM_PHOTO (#82)
If there is evidence of INDIUM_PRESENT
And SPECTRA.TYPE is "PHOTO"
Then INDIUM_ANALYZED_PHOTO is confirmed.
And CODE is set to "IN"
And 0.0 is assigned to INDIUM_CALIBRATION_PEAK.AREA
RULE : Rule INDIUM_LINE_X4 (#83)
If there is evidence of INDIUM_PRESENT
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is greater than 872
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is less than 874
Then INDIUM_LINE
is confirmed.
And IN_LINE_X4.EXIST is set to TRUE
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is assigned to IN_LINE_X4.WN

RULE : Rule INDIUM_LINE_X3 (#84)
If there is evidence of INDIUM_PRESENT
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is greater than 862
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is less than 864
Then INDIUM_LINE
is confirmed.
And IN_LINE_X3.EXIST is set to TRUE
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is assigned to IN_LINE_X3.WN

RULE : Rule INDIUM_LINE_X2 (#85)
If there is evidence of INDIUM_PRESENT
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is greater than 830
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is less than 832
Then INDIUM_LINE
is confirmed.
And IN_LINE_X2.EXIST is set to TRUE
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is assigned to IN_LINE_X2.WN
RULE : Rule INDIUM_LINE_X1 (#86)
    If
        there is evidence of INDIUM_PRESENT
        And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is greater than 7.98
        And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is less than 780
    Then INDIUM_LINE
        is confirmed.
        And IN_LINE_X1.EXIST is set to TRUE
        And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is assigned to IN_LINE_X1.WN

RULE : Rule INDIUM_LINE_EX (#87)
    If
        there is evidence of INDIUM_PRESENT
        And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is greater than 908
        And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is less than 910
    Then INDIUM_LINE
        is confirmed.
        And IN_LINE_EX.EXIST is set to TRUE
        And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is assigned to IN_LINE_EX.WN

RULE : Rule INDIUM_LINE_ESTAR (#88)
    If
        there is evidence of INDIUM_PRESENT
        And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is greater than 1609.2
        And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is less than 1610.2
    Then INDIUM_LINE
        is confirmed.
        And IN_LINE_ESTAR.EXIST is set to TRUE
        And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is assigned to IN_LINE_ESTAR.WN

103
RULE : Rule INDIUM_LINE_El (#89)
If
there is evidence of INDIUM_PRESENT
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is greater than 1254.3
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is less than 1255.3
Then INDIUM_LINE
is confirmed.
And IN_LINE_El.EXIST is set to TRUE
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is assigned to IN_LINE_El.WN

RULE : Rule INDIUM_LINE_9 (#90)
If
there is evidence of INDIUM_PRESENT
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is greater than 1245.3
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is less than 1246.3
Then INDIUM_LINE
is confirmed.
And IN_LINE_9.EXIST is set to TRUE
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is assigned to IN_LINE_9.WN

RULE : Rule INDIUM_LINE_8 (#91)
If
there is evidence of INDIUM_PRESENT
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is greater than 1241
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is less than 1244
Then INDIUM_LINE
is confirmed.
And IN_LINE_8.EXIST is set to TRUE
And <|POSSIBLE_INDIUM_PEAKS|>.WAVENUMBER is assigned to IN_LINE_8.WN
RULE : Rule INDIUM_LINE_7 (#92)
If
  there is evidence of INDIUM_PRESENT
  And <POSSIBLE_INDIUM_PEAKS>.WAVENUMBER is greater than
    1238.5
  And <POSSIBLE_INDIUM_PEAKS>.WAVENUMBER is less than
    1239.5
Then INDIUM_LINE
  is confirmed.
  And IN_LINE_7.EXIST is set to TRUE
  And <POSSIBLE_INDIUM_PEAKS>.WAVENUMBER is assigned to
    IN_LINE_7.WN

RULE : Rule INDIUM_LINE_6 (#93)
If
  there is evidence of INDIUM_PRESENT
  And <POSSIBLE_INDIUM_PEAKS>.WAVENUMBER is greater than
    1236
  And <POSSIBLE_INDIUM_PEAKS>.WAVENUMBER is less than
    1237
Then INDIUM_LINE
  is confirmed.
  And IN_LINE_6.EXIST is set to TRUE
  And <POSSIBLE_INDIUM_PEAKS>.WAVENUMBER is assigned to
    IN_LINE_6.WN

RULE : Rule INDIUM_LINE_5 (#94)
If
  there is evidence of INDIUM_PRESENT
  And <POSSIBLE_INDIUM_PEAKS>.WAVENUMBER is greater than
    1231.7
  And <POSSIBLE_INDIUM_PEAKS>.WAVENUMBER is less than
    1232.7
Then INDIUM_LINE
  is confirmed.
  And IN_LINE_5.EXIST is set to TRUE
  And <POSSIBLE_INDIUM_PEAKS>.WAVENUMBER is assigned to
    IN_LINE_5.WN

105
RULE : Rule INDIUM_LINE_4P (#95)
If
there is evidence of INDIUM_PRESENT
And <\POSSIBLE_INDIUM PEAKS>.WAVENUMBER is greater than 1598
And <\POSSIBLE_INDIUM PEAKS>.WAVENUMBER is less than 1599
Then INDIUM_LINE
is confirmed.
And IN_LINE_4P.EXIST is set to TRUE
And <\POSSIBLE_INDIUM PEAKS>.WAVENUMBER is assigned to IN_LINE_4P.WN

RULE : Rule INDIUM_LINE_4A (#96)
If
there is evidence of INDIUM_PRESENT
And <\POSSIBLE_INDIUM PEAKS>.WAVENUMBER is greater than 1217.5
And <\POSSIBLE_INDIUM PEAKS>.WAVENUMBER is less than 1218.5
Then INDIUM_LINE
is confirmed.
And IN_LINE_4A.EXIST is set to TRUE
And <\POSSIBLE_INDIUM PEAKS>.WAVENUMBER is assigned to IN_LINE_4A.WN

RULE : Rule INDIUM_LINE_4 (#97)
If
there is evidence of INDIUM_PRESENT
And <\POSSIBLE_INDIUM PEAKS>.WAVENUMBER is greater than 1215.3
And <\POSSIBLE_INDIUM PEAKS>.WAVENUMBER is less than 1217.3
Then INDIUM_LINE
is confirmed.
And IN_LINE_4.EXIST is set to TRUE
And <\POSSIBLE_INDIUM PEAKS>.WAVENUMBER is assigned to IN_LINE_4.WN

106
RULE : Rule INDIUM_LINE_3P (#98)
If there is evidence of INDIUM_PRESENT
And <$\text{POSSIBLE_INDIUM_PEAKS}$>.WAVENUMBER is greater than 1589.3
And <$\text{POSSIBLE_INDIUM_PEAKS}$>.WAVENUMBER is less than 1590.3
Then INDIUM_LINE
is confirmed.
And IN_LINE_3P.EXIST is set to TRUE
And <$\text{POSSIBLE_INDIUM_PEAKS}$>.WAVENUMBER is assigned to IN_LINE_3P.WN

RULE : Rule INDIUM_LINE_3 (#99)
If there is evidence of INDIUM_PRESENT
And <$\text{POSSIBLE_INDIUM_PEAKS}$>.WAVENUMBER is greater than 1207.5
And <$\text{POSSIBLE_INDIUM_PEAKS}$>.WAVENUMBER is less than 1208.5
Then INDIUM_LINE
is confirmed.
And IN_LINE_3.EXIST is set to TRUE
And <$\text{POSSIBLE_INDIUM_PEAKS}$>.WAVENUMBER is assigned to IN_LINE_3.WN

RULE : Rule INDIUM_LINE_2P (#100)
If there is evidence of INDIUM_PRESENT
And <$\text{POSSIBLE_INDIUM_PEAKS}$>.WAVENUMBER is greater than 1565
And <$\text{POSSIBLE_INDIUM_PEAKS}$>.WAVENUMBER is less than 1567
Then INDIUM_LINE
is confirmed.
And IN_LINE_2P.EXIST is set to TRUE
And <$\text{POSSIBLE_INDIUM_PEAKS}$>.WAVENUMBER is assigned to IN_LINE_2P.WN
RULE: Rule INDIUM_LINE_2 (#101)
If
there is evidence of INDIUM_PRESENT
And <[POSSIBLE_INDIUM_PEAKS]> WAVENUMBER is greater than 1175.3
And <[POSSIBLE_INDIUM_PEAKS]> WAVENUMBER is less than 1176.3
Then INDIUM_LINE
is confirmed.
And IN_LINE_2.EXIST is set to TRUE
And <[POSSIBLE_INDIUM_PEAKS]> WAVENUMBER is assigned to IN_LINE_2.WN

RULE: Rule INDIUM_LINE_11 (#102)
If
there is evidence of INDIUM_PRESENT
And <[POSSIBLE_INDIUM_PEAKS]> WAVENUMBER is greater than 1253
And <[POSSIBLE_INDIUM_PEAKS]> WAVENUMBER is less than 1255
Then INDIUM_LINE
is confirmed.
And IN_LINE_11.EXIST is set to TRUE
And <[POSSIBLE_INDIUM_PEAKS]> WAVENUMBER is assigned to IN_LINE_11.WN

RULE: Rule INDIUM_LINE_10 (#103)
If
there is evidence of INDIUM_PRESENT
And <[POSSIBLE_INDIUM_PEAKS]> WAVENUMBER is greater than 1247.4
And <[POSSIBLE_INDIUM_PEAKS]> WAVENUMBER is less than 1251.4
Then INDIUM_LINE
is confirmed.
And IN_LINE_10.EXIST is set to TRUE
And <[POSSIBLE_INDIUM_PEAKS]> WAVENUMBER is assigned to IN_LINE_10.WN
RULE : Rule INDIUM_LINE_1 (#104)
If
there is evidence of INDIUM_PRESENT
And <POSIBLE_INDIUM_PEAKS>.WAVENUMBER is greater than 1145
And <POSIBLE_INDIUM_PEAKS>.WAVENUMBER is less than 1146
Then INDIUM_LINE
is confirmed.
And IN_LINE_1.EXIST is set to TRUE
And <POSIBLE_INDIUM_PEAKS>.WAVENUMBER is assigned to IN_LINE_1.WN

RULE : Rule INVESTIGATE_INDIUM (#105)
If
there is evidence of POSSIBLE_INDIUM
And <POSIBLE_INDIUM_PEAKS>.WAVENUMBER is greater than 1144.4
And <POSIBLE_INDIUM_PEAKS>.WAVENUMBER is less than 1146.4
And <<POSIBLE_INDIUM_PEAKS>>.WAVENUMBER is greater than 1174.8
And <<POSIBLE_INDIUM_PEAKS>>.WAVENUMBER is less than 1176.8
And <<POSIBLE_INDIUM_PEAKS>>.WAVENUMBER is greater than 1215.3
And <<POSIBLE_INDIUM_PEAKS>>.WAVENUMBER is less than 1217.3
Then INDIUM_PRESENT
is confirmed.
And <<POSIBLE_INDIUM_PEAKS>>.WAVENUMBER is assigned to INDIUM_CALIBRATION_PEAK.WAVENUMBER
RULE: Rule INVESTIGATE_OXYGEN_ABS (#106)
If
    there is evidence of OXYGEN_PRESENT
    And SPECTRA.TYPE is "ABS"
Then OXYGEN_ANALYZED_ABS is confirmed.
    And CODE is set to "OX"
    And Execute "FINDAREA"(@ATOMID=SPECTRA.POINTS,OXYGEN_CALIBRATION_PEAK.WAVENUMBER,CODE.Value;)
    And (OXYGEN_CALIBRATION_PEAK.AREA*3.090e+16) is assigned to OXYGEN_CALIBRATION_PEAK.CONC

RULE: Rule INVESTIGATE_OXYGEN_PHOTO (#107)
If
    there is evidence of OXYGEN_PRESENT
    And SPECTRA.TYPE is "PHOTO"
Then OXYGEN_ANALYZED_PHOTO is confirmed.
    And CODE is set to "OX"
    And 0.0 is assigned to OXYGEN_CALIBRATION_PEAK.AREA

RULE: Rule OXYGEN_LINE_1 (#108)
If
    there is evidence of OXYGEN_PRESENT
    And <POSIBLE_OXYGEN_PEAKS>.WAVENUMBER is greater than 1135.0
    And <POSIBLE_OXYGEN_PEAKS>.WAVENUMBER is less than 1137.0
Then OXYGEN_LINE is confirmed.
    And OX_LINE_1.EXIST is set to TRUE
    And <POSIBLE_OXYGEN_PEAKS>.WAVENUMBER is assigned to OX_LINE_1.WN
RULE : Rule INVESTIGATE_OXYGEN (#109)
If
there is evidence of POSSIBLE_OXYGEN
And <PPOSSIBLE_OXYGEN_PEAKS>.WAVENUMBER is greater than 1135.0
And <PPOSSIBLE_OXYGEN_PEAKS>.WAVENUMBER is less than 1137.0
Then OXYGEN_PRESENT
is confirmed.
And <PPOSSIBLE_OXYGEN_PEAKS>.WAVENUMBER is assigned to OXYGEN_CALIBRATION_PEAK.WAVENUMBER

RULE : Rule EXAMINE_THE_POSSIBLE_PEAKS (#110)
If
GALLIUM_ANALYZED_ABS is not NOTKNOWN
And GALLIUM_ANALYZED_PHOTO is not NOTKNOWN
And BORON_ANALYZED_ABS is not NOTKNOWN
And BORON_ANALYZED_PHOTO is not NOTKNOWN
And INDIUM_ANALYZED_ABS is not NOTKNOWN
And INDIUM_ANALYZED_PHOTO is not NOTKNOWN
And ALUMINUM_ANALYZED_ABS is not NOTKNOWN
And ALUMINUM_ANALYZED_PHOTO is not NOTKNOWN
And CARBON_ANALYZED_ABS is not NOTKNOWN
And CARBON_ANALYZED_PHOTO is not NOTKNOWN
And OXYGEN_ANALYZED_ABS is not NOTKNOWN
And OXYGEN_ANALYZED_PHOTO is not NOTKNOWN
Then PEAKS_ANALYZED
is confirmed.

RULE : Rule Force_the_impurity_analysis (#111)
If
POSSIBLE_ALUMINUM is not NOTKNOWN
And POSSIBLE_GALLIUM is not NOTKNOWN
And POSSIBLE_INDIUM is not NOTKNOWN
And POSSIBLE_BORON is not NOTKNOWN
And POSSIBLE_CARBO is not NOTKNOWN
And POSSIBLE_OXYGEN is not NOTKNOWN
And POSSIBLE_UNKNOWN_PEAKS is not NOTKNOWN
Then PEAKS_EXAMINED
is confirmed.
RULE: Rule Load_the_peak_table (#112)

If

there is evidence of SPECTRA_ID ASKED
And SPECTRA_TYPE is "ABS", "PHOTO"

Then PEAKS_LOADED
is confirmed.
And Retrieve from "@V(SPECTRA.PEAKS)"
@TYPE=NXPDB; @FILL=ADD; @UNKNOWN=TRUE; @NAME="NAME";
@CREATE=POSSIBLE_PEAKS; @PROPS=AMPLITUDE, WAVENUMBER,
HALF_WIDTH, AREA; @FIELDS="AMPLITUDE", "WAVENUMBER",
"HALF_WIDTH", "AREA";

RULE: Rule PICK_POSSIBLE_ALUMINUM_PEAKS (#113)

If

<|POSSIBLE_PEAKS|>. WAVENUMBER is greater than 337
And <|POSSIBLE_PEAKS|>. WAVENUMBER is less than 912

Then POSSIBLE_ALUMINUM
is confirmed.
And Create Object <|POSSIBLE_PEAKS|>
|POSSIBLE_ALUMINUM_PEAKS|

RULE: Rule PICK_OUT_POSSIBLE_BORON_PEAKS (#114)

If

<|POSSIBLE_PEAKS|>. WAVENUMBER is greater than 183
And <|POSSIBLE_PEAKS|>. WAVENUMBER is less than 713.5

Then POSSIBLE_BORON
is confirmed.
And Create Object <|POSSIBLE_PEAKS|>
|POSSIBLE_BORON_PEAKS|

RULE: Rule PICK_POSSIBLE_CARBON_PEAK (#115)

If

<|POSSIBLE_PEAKS|>. WAVENUMBER is greater than 599.0
And <|POSSIBLE_PEAKS|>. WAVENUMBER is less than 601.0

Then POSSIBLE_CARBON
is confirmed.
And Create Object <|POSSIBLE_PEAKS|>
|POSSIBLE_CARBON_PEAKS|
RULE : Rule PICK_POSSIBLE_GALLIUM_PEAKS (#116)
   If
   
   <POSSIBLE_PEAKS>.WAVENUMBER is greater than 346
   And <POSSIBLE_PEAKS>.WAVENUMBER is less than 942
   Then POSSIBLE_GALLIUM
   is confirmed.
   And Create Object <POSSIBLE_PEAKS>
   |POSSIBLE_GALLIUM_PEAKS|

RULE : Rule PICK_OUT_POSSIBLE_INDIUM_PEAKS (#117)
   If
   
   <POSSIBLE_PEAKS>.WAVENUMBER is greater than 798
   And <POSSIBLE_PEAKS>.WAVENUMBER is less than 1610
   Then POSSIBLE_INDIUM
   is confirmed.
   And Create Object <POSSIBLE_PEAKS>
   |POSSIBLE_INDIUM_PEAKS|

RULE : Rule PICK_POSSIBLE_OXYGEN_PEAKS (#118)
   If
   
   <POSSIBLE_PEAKS>.WAVENUMBER is greater than 1135.0
   And <POSSIBLE_PEAKS>.WAVENUMBER is less than 1137.0
   Then POSSIBLE_OXYGEN
   is confirmed.
   And Create Object <POSSIBLE_PEAKS>
   |POSSIBLE_OXYGEN_PEAKS|
RULE : Rule UNIDENTIFIED_PEAKS (#119)

If
<br/>&lt;POSSIBLE_PEAKS|&gt; is not a member of
<br/>&lt;POSSIBLE_ALUMINUM_PEAKS|&gt;
And &lt;POSSIBLE_PEAKS|&gt; is not a member of
&lt;POSSIBLE_BORON_PEAKS|&gt;
And &lt;POSSIBLE_PEAKS|&gt; is not a member of
&lt;POSSIBLE_INDIUM_PEAKS|&gt;
And &lt;POSSIBLE_PEAKS|&gt; is not a member of
&lt;POSSIBLE_GALLIUM_PEAKS|&gt;
And &lt;POSSIBLE_PEAKS|&gt; is not a member of
&lt;POSSIBLE_CARBON_PEAKS|&gt;
And &lt;POSSIBLE_PEAKS|&gt; is not a member of
&lt;POSSIBLE_OXYGEN_PEAKS|&gt;
Then POSSIBLE_UNKNOWN_PEAKS
is confirmed.
And Create Object &lt;POSSIBLE_PEAKS|&gt; |UNIDENTIFIED_PEAKS|

RULE : Rule PLOT_THE_SPECTRA (#120)

If
<br/>REPORT_ALUMINUM is not NOTKNOWN
And REPORT_BORON is not NOTKNOWN
And REPORT_GALLIUM is not NOTKNOWN
And REPORT_INDIUM is not NOTKNOWN
And REPORT_OXYGEN is not NOTKNOWN
And REPORT_CARBON is not NOTKNOWN
Then reports_done
is confirmed.
And Execute "rgraph"(@ATOMID=SPECTRA.NAME;)

RULE : Rule Get_the_spectra_ID (#121)

If
<br/>SPECTRA.NAME is assigned to SPECTRA.NAME
Then SPECTRA_ID_ASKED
is confirmed.
And Execute "FILLSPECTRA"(@ATOMID=SPECTRA.NAME;)
And Execute "PEAKPICK"()
And STRCAT(SPECTRA.NAME,".MAS") is assigned to SPECTRA.POINTS
And STRCAT(SPECTRA.NAME,".NXP") is assigned to SPECTRA.PEAKS