(54) Title: DEHYDRATION AND PURIFICATION OF ISOPROPYL ALCOHOL

(57) Abstract

A method and apparatus for the on-site reprocessing of isopropyl alcohol used in semiconductor manufacturing, to generate an ultrapure and ultratray isopropyl alcohol, produced through a pervaporation step, followed by a two-step distillation, wherein the first distillation step, an autonomous azeotropic self-stripping distillation column (78) is used to produce an ultradry and partially purified isopropyl alcohol, and in the next distillation step, the isopropyl alcohol is distilled in an overhead product distillation column (164), to produce an ultrapure and ultradry isopropyl alcohol.
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DEHYDRATION AND PURIFICATION OF ISOPROPYL ALCOHOL

Background of the Invention

This invention relates generally to an apparatus and method for purification and dehydration of isopropyl alcohol. More particularly, the present invention relates to a method and apparatus suitable for location at a semiconductor manufacturing facility and useful for continuous on-site dehydration and purification of isopropyl alcohol to ultradry and ultrapure levels.

Many steps in the semiconductor wafer manufacturing process are followed by a deionized water rinse, which is then followed by a drying step. During this wafer drying step, it is important to prevent watermarks from forming on the surface of the silicon wafers. Watermarks typically form when silicon dioxide and other dissolved contaminants precipitate out of the deionized water as it evaporates from the surface of the wafer. The presence of watermarks on a partially manufactured wafer creates serious difficulties in subsequent manufacturing processes.

Watermark formation on silicon wafers can be minimized or prevented by keeping the deionized water from evaporating off of the wafer surface during the drying process. Several important techniques for achieving this result involve the use of isopropyl alcohol (IPA). In one such technique, the water on the surface of the wafer is displaced by isopropyl alcohol before the water has a chance to evaporate, and then the alcohol is evaporated from the surface of the wafer. Another technique, called vapor drying, involves condensation of isopropyl alcohol vapor onto the surface of the wafer, causing the water present on the wafer to be taken up by the dry alcohol. The water-rich alcohol then drips off of the wafer before water evaporation can occur, and is replaced by more dry alcohol condensate, which is then evaporated.

To minimize or prevent watermarks and to enhance drying, semiconductor manufacturers prefer using ultrapure and ultradry isopropyl alcohol. Ultrapure is defined here as having zero particles per milliliter of a size larger
than 2.0 microns, having zero to 2 particles per milliliter of a size of 0.5 microns to 2.0 microns, having zero to 30 particles per milliliter of a size between 0.1 microns and 0.5 microns, having an unspecified number of particles per milliliter below 0.1 microns, having 1 part per trillion (ppt) to 1 part per billion (ppb) of any specific trace impurity such as metals, anions, and cations, and having 10 ppt to 10 parts per million (ppm) of any other specific trace organic substances. Ultradry is defined here as having between a high of 100 parts per million (ppm) and a low of 0.1 ppm of water in the isopropyl alcohol.

Currently, the availability of ultradry and ultrapure isopropyl alcohol from suppliers is limited in relation to the demands of the industry for the chemical. In addition, ultrapure and ultradry isopropyl alcohol purchased from off-site suppliers may lose its purity due to contaminants or water added during its handling and transportation to the semiconductor manufacturer. Moreover, transporting thousands of liters of hazardous chemicals over public highways every month is undesirable.

Even if a semiconductor manufacturer is fortunate enough to secure an adequate supply of ultrapure and ultradry isopropyl alcohol, once the chemical is used to dry semiconductor wafers and picks up contamination and water, it is no longer useful as a drying agent and must be disposed of. In the past, this has involved resale to industries demanding less pure isopropyl alcohol, or disposal as a hazardous chemical, or sending the isopropyl alcohol off-site for recycling. All of these disposal methods create additional expenses for the semiconductor manufacturer, which are passed on to consumers in the form of higher prices.

The best solution to these problems is for the semiconductor manufacturer to recycle and purify the used isopropyl alcohol to an ultrapure and ultradry level at the manufacturing facility. Unfortunately, the current methods of purifying and dehydrating isopropyl alcohol are not suited to meet this need. For example, one well-known method of purifying isopropyl alcohol involves simple
overhead product distillation. This method, while useful in removing contaminants with boiling points higher than isopropyl alcohol, cannot be used to dehydrate isopropyl alcohol to an ultradry level, because isopropyl alcohol forms a low boiling azeotrope with water. In addition, this method also does nothing to remove those contaminants with boiling points lower than isopropyl alcohol.

Another method used to dehydrate isopropyl alcohol involves use of pervaporation membranes. Pervaporation membranes can be used to dry isopropyl alcohol with a high initial water content to water levels around 300 to 500 ppm. It is extremely difficult, however, to use pervaporation to dehydrate isopropyl alcohol to an ultradry level having a water content of less than 100 ppm. Moreover, depending upon the type of pervaporation membrane used, pervaporation may actually add particles and organic materials to the isopropyl alcohol. Thus, pervaporation, by itself, cannot be used to recycle used isopropyl alcohol to an ultrapure and ultradry level.

Another method teaches the use of a molecular sieve to dehydrate the isopropyl alcohol, followed by overhead product distillation for purification. The combination of the molecular sieve process and distillation, however, does nothing to remove most light organic substances or other contaminants with low boiling points. Thus, while this method can work better than previous methods in achieving ultradry isopropyl alcohol, contaminants with boiling points lower than isopropyl alcohol may still remain after treatment.

Another prior art method teaches the process of adding a second liquid to an alcohol-water mix, such as propane as described in U.S. Patent No. 5,053,563, or di-isopropyl ether as described in U.S. Patent No. 4,762,616. The second liquid is added because it forms a lower boiling point azeotrope with water than the azeotrope of isopropyl alcohol and water. Two column distillation is then used to break the isopropyl alcohol-water azeotrope and dehydrate the alcohol. While this process might work for isopropyl alcohol, the complexity of the process makes it difficult
for use as an on-site recycling process for use at semiconductor manufacturing sites. In addition, the use of a second liquid raises serious questions regarding the possible introduction of additional impurities.

For the foregoing reasons, there is a need for a method and apparatus, suitable for location at a semiconductor manufacturing facility, which can be used to recycle isopropyl alcohol to ultradry and ultrapure levels at the cost and volumes needed by the semiconductor manufacturing industry.

**Summary of the Invention**

The present invention is directed to a method and apparatus that satisfies the need of the semiconductor manufacturer for dehydrating and purifying used isopropyl alcohol to an ultradry and ultrapure level.

In accordance with one aspect of the present invention, there is provided a method for dehydrating and purifying a solution containing isopropyl alcohol, water, and minor amounts of other contaminants to produce an ultradry and ultrapure isopropyl alcohol. The first step of this method involves removing water from a solution containing isopropyl alcohol by pervaporation of the solution through a water-permeable membrane. This produces a partially dehydrated isopropyl alcohol solution which contains a small amount of water, and minor amounts of other contaminants. The next step is to distill the partially dehydrated isopropyl alcohol in an autonomous azeotropic self-stripping distillation column, to remove substantially all of the remaining water and low boiling point contaminants, thereby producing an ultradry and partially purified isopropyl alcohol which is removed from a reboiler at the column bottom. The next step in the method involves distilling the ultradry and partially purified isopropyl alcohol through a low-boiling overhead product distillation column, then taking the ultradry and ultrapure isopropyl alcohol as an end product overhead condensate.

The advantages of the method of the present invention are that it provides a relatively simple process to recycle
used isopropyl alcohol to an ultrapure and ultradry level, allowing less complicated and lower cost equipment to be installed at the semiconductor manufacturing facility. Furthermore, the method of the present invention is adapted to work with isopropyl alcohol solutions having wide variations in water content, so that even if the isopropyl alcohol solution is severely contaminated with water, the present method can be used to dehydrate and purify it. The present method also avoids the need for the addition of a second chemical to break the isopropyl alcohol-water azeotrope.

In accordance with another aspect of the present invention, there is provided a method of dehydrating and purifying an isopropyl alcohol solution containing only minor amounts of organic and inorganic contaminants, and less than about 2,000 ppm water, to an ultrapure and ultradry level. In this embodiment, the solution to be dehydrated and purified is not passed through a pervaporation membrane, as discussed previously, but is instead first introduced into an autonomous azeotropic self-stripping distillation column. This first distillation step removes substantially all of the water and low boiling point contaminants from the isopropyl alcohol solution, to produce an ultradry and partially purified isopropyl alcohol which is removed from a reboiler at the column bottom. The next step involves distilling the ultradry and partially purified isopropyl alcohol through a low-boiling overhead product distillation column, then taking the ultradry and ultrapure isopropyl alcohol as an end product overhead condensate.

Preferably, in each described distillation step used in practicing the methods of the present invention, the distillation column will have from 11 to 40 theoretical distillation plates. In addition, in the preferred methods of the present invention, the ultrapure and ultradry isopropyl alcohol end product is subsequently filtered to remove submicron sized particulate impurities.

In another aspect of the present invention, there is provided an apparatus which can be used to practice the
above-described methods. An apparatus having the features of the present invention comprises an isopropyl alcohol input line which is connected to a pervaporation chamber, or to other means for pervaporation of an isopropyl alcohol solution. An output line from the pervaporation chamber is attached to a first distillation system, which comprises a preheating column connected to an autonomous azeotropic self-stripping distillation column, where the distillation column includes a condenser at its top, and a reboiler at its bottom. The reboiler of the distillation column has an output line which is adapted to permit removal of an ultradry and partially purified isopropyl alcohol product after the distillation process has occurred. The output from this reboiler is connected to a second distillation system which comprises a low boiling overhead product distillation column. This second distillation column has a reboiler at its bottom, and an end product condenser at its top. A product cooling column connected to the end product condenser is used to cool the distilled ultrapure and ultradry isopropyl alcohol after the distillation, and the end product then passes into an end product output line connected to the cooling column.

In accordance with another aspect of the present invention, there is provided an apparatus useful for dehydrating and purifying isopropyl alcohol solutions containing less than about 2000 ppm water. The apparatus of this embodiment features an isopropyl alcohol input line which is attached to a first distillation system. The first distillation system comprises a preheating column connected to an autonomous azeotropic self-stripping distillation column, where the distillation column includes a condenser at its top, and a reboiler at its bottom. The reboiler of the distillation column has an output line adapted to permit removal of an ultradry and partially purified isopropyl alcohol product after the distillation process has occurred. The output from this reboiler is connected to a second distillation system which comprises a low boiling overhead product distillation column. This
second distillation column has a reboiler at its bottom, and an end product condenser at its top. A product cooling column connected to the end product condenser is used to cool the distilled ultrapure and ultradry isopropyl alcohol after the distillation, and the end product then passes into an end product output line connected to the cooling column.

Preferably, the distillation columns of the preferred embodiments of the present invention are constructed in such a manner that they comprise 11 to 40 theoretical plates. In addition, in the preferred embodiments of the present invention, filtration means are connected to the end product output line to remove any submicron sized particulate impurities passed through the filtration means.

These and other features and advantages of the present invention will become apparent from the detailed description of the preferred embodiments which follows, when taken together with the attached drawings and claims.

**Brief Description of the Drawings**

Fig. 1 shows a block diagram of the preferred apparatus used in accordance with the method of the present invention.

Fig. 2 is a schematic diagram of the preferred apparatus used to practice the pervaporation step of the invention.

Fig. 3 is a schematic diagram of the preferred apparatus used to practice the first distillation step of the method of the present invention.

Fig. 4 is a schematic diagram of the preferred apparatus used to practice the second distillation step of the method of the present invention.

Fig. 5a shows the process flow diagram of the preferred apparatus used in accordance with the method and process of the present invention for the conditions given in example 1. Fig. 5b shows the process flow diagram of the preferred apparatus used in accordance with the method and process of the present invention for the conditions given in example 2.
Fig. 6 shows a set of control rules which may be executed by hardware and software controllers which can be used in the present invention.

**Detailed Description of Preferred Embodiments**

In one preferred embodiment, the method according to the present invention for dehydrating and purifying isopropyl alcohol to an ultradry and ultrapure level begins with pervaporation of a solution containing isopropyl alcohol, water, and minor amounts of various other contaminants, thereby producing an impure and partially dehydrated isopropyl alcohol containing about 500 ppm water. The isopropyl alcohol from the pervaporation step is then channeled to a first distillation step, where the partially dehydrated isopropyl alcohol is distilled in an autonomous azeotropic self-stripping distillation column to remove substantially all of the remaining water and other low-boiling contaminants, thereby forming an ultradry isopropyl alcohol. The ultradry isopropyl alcohol from the first distillation step is then channeled to a second distillation step, to remove impurities with boiling points higher than isopropyl alcohol, thereby forming an ultrapure and ultradry isopropyl alcohol product. Optionally, a filtration step may be added after the second distillation step to further purify the ultrapure and ultradry isopropyl alcohol.

In another preferred embodiment of the present invention, applicable when the isopropyl alcohol to be purified and dehydrated contains less than 2000 ppm water, no pervaporation step is used in the dehydration process. In this embodiment, the isopropyl alcohol solution to be purified and dehydrated is directly channeled to the first distillation step where the isopropyl alcohol is distilled in an autonomous azeotropic self-stripping distillation column to remove substantially all of the water and other low-boiling contaminants, thus forming an ultradry alcohol. The ultradry alcohol from the first distillation step is then channeled to a second distillation step, to remove impurities with boiling points higher than isopropyl alcohol, thereby forming an ultrapure and ultradry alcohol.
isopropyl alcohol product. Optionally, a filtration step may be added after the second distillation step to further purify the ultrapure and ultradry isopropyl alcohol.

Another preferred embodiment of the present invention is an apparatus, small enough to be located at a semiconductor manufacturing facility, and capable of practicing the methods of the present invention to purify and dehydrate isopropyl alcohol to an ultrapure and ultradry level.

Figures 1 and 2 disclose a representation of a preferred embodiment of an apparatus suitable for carrying out the pervaporation step of the present invention. The pervaporation step of the present invention removes the excess water from the isopropyl alcohol, producing a partially dehydrated isopropyl alcohol containing about 500 ppm water. Pervaporation is especially advantageous when the isopropyl alcohol to be dehydrated contains water in excess of the azeotropic concentration, as conventional distillation is ineffective in separating the water-isopropyl alcohol azeotrope.

Referring now to Figure 2, the pervaporation sub-system 600 comprises a pre-heater column 614, a pervaporation cell 620, a pervaporation vacuum apparatus 668, and a pervaporation waste water condenser 618. Isopropyl alcohol to be processed, containing from 0.2% to 12% by weight water, plus from 1 ppm to 3% other impurities, enters the pervaporation feed line 510, passes through metering pump 650, and flows into the pervaporation pre-heater column 614. Preferably, pre-heater column 614 heats the isopropyl alcohol to a temperature of about 50 degrees Celsius, thereby optimizing the removal of water during the subsequent pervaporation. Heat is supplied to the pre-heater column 614 from a steam source line 90 through a feed line 626. The steam flow through pre-heater column 614 is controlled by an inlet steam control valve 628, thereby permitting the regulation of the temperature of pre-heater column 614. The resulting steam condensate passing through pre-heater column 614 exits through a condensate return line 624 to a steam return line 92, for
recycling through the steam subsystem 400. After being heated within the pre-heater column 614, the isopropyl alcohol flows through a pervaporation cell feed line 676 and into the pervaporation cell 620. Preferably, a thermocouple 630 is positioned on the feed line 676 to facilitate temperature measurements of the heated isopropyl alcohol before it is input into the pervaporation cell 620.

Pervaporation cell 620 comprises a pervaporation dehydration chamber 692, a pervaporation vaporetate chamber 652, and a water permeable pervaporation membrane 654, where the pervaporation membrane 654 separates the pervaporation dehydration chamber 692 from the pervaporation vaporetate chamber 652. In the most preferred embodiment, the pervaporation membrane 654 is highly permeable to water, but substantially impermeable to isopropyl alcohol. Advantageously, the atmospheric pressure of pervaporation chamber 652 is reduced relative to the pervaporation dehydration chamber 692 to facilitate diffusion of water across pervaporation membrane 654, and its subsequent evaporation in the pervaporation vaporetate chamber 652. In the illustrated embodiment, this is achieved by vacuum apparatus 668 which is attached to the pervaporation vaporetate chamber 652 by vacuum line 678. In practice, the pervaporation cell 620 removes water from isopropyl alcohol because the pervaporation membrane 654 is both permeable to water and has a greater chemical affinity for water than for isopropyl alcohol. Thus, water molecules entering into the pervaporation dehydration chamber 692 adhere to the pervaporation membrane 654, then diffuse across the membrane and evaporate from the membrane into pervaporation vaporetate chamber 652.

Water vapor passing through the pervaporation membrane 654 condenses in the pervaporation vaporetate chamber 692, and is subsequently drawn into vacuum line 678 to be deposited in a waste water condenser 618. Preferably, a vacuum sensor 666 is positioned within the pervaporation vaporetate chamber 692 to facilitate measurement of the level of vacuum created by the vacuum apparatus 668. The water in the waste water condenser 618
is removed from the pervaporation sub-system by a metering pump 682, and is delivered to a waste drain 694 through the waste water line 610. Cooling water enters into the waste water condenser 618 through a cooling water input line 662, flows through condenser coil 656, and exits through a cooling water output line 660. The flow of cooling water through condenser coil 656 is controlled by a water control valve 658. Optionally, a thermocouple 636 can be positioned within the waste water condenser 618 to facilitate measurement of the waste water temperature. Also optionally, a level sensor 638 can be positioned within pervaporation waste water condenser 618 to facilitate measurement of the waste water level. In the embodiment illustrated in Figure 2, an exhaust 690 is required for proper operation.

Isopropyl alcohol which is partially dehydrated in the pervaporation dehydration chamber 692 flows from the pervaporation cell 620 through a pervaporation sub-system output line 686 to distillation subsystem 10 for further processing. Optionally, in a more preferred embodiment, a water content monitor 632 is positioned within the pervaporation dehydration chamber to facilitate the measurement of the isopropyl alcohol water content. The water content monitor 632 may be an automatic concentration gauge based upon the dielectric constant of the isopropyl alcohol, or it may be an automatic Karl Fisher Titrator that determines the water content directly, or any other concentration monitoring means one skilled in the art may employ.

Pervaporation units suitable for practicing the present invention are well known in the art, and may be easily manufactured or obtained from a variety of suppliers. For example, a complete pervaporation unit suitable for practicing the present invention, called the MiniPervap plant, may be purchased from Carbone of America, Parsippany, New Jersey.

Figures 1 and 3 illustrate the details of a preferred embodiment of an apparatus suitable for carrying out the first distillation step of the present invention. The
first distillation step removes water remaining after the pervaporation step, bringing the isopropyl alcohol to an ultradry level of between 0.1 ppm to 100 ppm water. Alternately, if the starting isopropyl alcohol contains less than 2000 ppm water, the pervaporation step may be bypassed, and the isopropyl alcohol may be introduced directly into the first distillation step.

Referring to Figure 3, the first distillation sub-system 10 comprises a pre-heater column 14, an autonomous azeotropic self-stripping distillation column 78, a reboiler 18, a condenser 16, a waste product cooler 12, and a reboiler output line 68.

Isopropyl alcohol from the pervaporation step of the present invention, containing approximately 500 ppm of water and various other multi-ppm level contaminants, flows through a feed line 20 into the pre-heater column 14. Alternately, in an embodiment of the present invention bypassing the pervaporation step, isopropyl alcohol containing less than 2000 ppm water is input directly into feed line 20 where it flows to the pre-heater column 14. Preferably, the pre-heater 14 column heats the isopropyl alcohol to a temperature of about 82 degrees Celsius, thereby allowing the distillation process to begin immediately upon the introduction of the isopropyl alcohol into distillation column 78. Steam from the steam source line 90 passes through a steam feed line 26, and provides a heating source for the pre-heater column 14. The steam flow is controlled by the inlet steam control valve 28, and the resulting steam condensate exits through the condensate return line 24 to the steam return line 92, for recycling in the steam sub-system 400.

The heated isopropyl alcohol flows through the pre-heater column 14, into column feed line 76, and into distillation column 78. Preferably, a thermocouple 74 is positioned on the feed line 76 to facilitate temperature measurements of the isopropyl alcohol before it flows into the distillation column 78. The distillation column 78 uses high efficiency packing to provide from eleven to forty theoretical plates in the compact vertical space.
Preferably, distillation column 78 is packed such that it has twenty theoretical plates. More preferably, distillation column 78 is packed such that it has twenty-five theoretical plates. In the twenty-five plate embodiment, the isopropyl alcohol is introduced into the distillation column 78 preferably at a point corresponding to theoretical plate number fourteen out of twenty-five. The rectifying section 48 and the stripping section 54 of distillation column 78 provide for the separation of water and other low-boiling point contaminants from the isopropyl alcohol. Preferably, a differential pressure sensor 93 is positioned on a differential pressure column line 99, where column line 99 is connected to the reboiler 18 and the top and bottom of the distillation column 78, as illustrated in Figure 3, to facilitate measurement of the pressure across the distillation column 78.

The distillation column reboiler 18 provides a receptacle for fluids refluxing through the distillation column 78. Liquids in the reboiler 18 are heated by heat transfer from a reboiler steam coil 56. Steam coil 56 receives steam from a feed line 60, which in turn is connected to the steam source line 90. The flow of steam through steam coil 56 is controlled by a steam inlet control valve 58, which permits adjustment of the reboiler temperature by varying steam flow. Steam passing through the steam coil 56 condenses, and the resulting steam condensate flows through condensate return line 62 and into steam return line 92. Preferably, a differential pressure sensor 66 and a thermocouple 64 are positioned within the reboiler 18 to facilitate temperature and pressure measurements of the fluid within the reboiler. A reboiler output line 68 is provided in reboiler 18 to withdraw the ultradry isopropyl alcohol after the distillation process. Passing through reboiler outlet line 68, the ultradry isopropyl alcohol is sent to column output line 52 for delivery to the second distillation subsystem 100. A control valve 50 and feed flowmeter 80 are used to regulate the flow of the ultradry isopropyl alcohol into the second distillation sub-system 100.
Distillation column condenser 16 is situated on top of the distillation column 78. Cooling water from the water source line 86 flows into the condensing coil 44 within the column condenser 16, causing the condensation of gaseous vapors rising up through the distillation column 78, for proper reflux back into column rectifying section 48. The cooling water exits condensing coil 44, flowing through a condenser outlet water control valve 46 and into the water return line 88, eventually returning to a cooling water sub-system 410, which recycles cooling water for reuse system-wide. Preferably, a thermocouple 42 and a pressure sensor 40 are positioned within the condenser 16 to facilitate measurement of pressure and temperature. Also preferably, a differential pressure sensor 95 is positioned on an inter-column differential pressure line 98, where pressure line 98 connects distillation column 78 to distillation column 164, depicted in Figure 4, to provide pressure measurements between the condenser 16 and a second distillation column condenser 104, also shown in Figure 4.

The conduit piping for differential pressure sensor 97 is used to provide pressure commutation to the pressure gauges.

Distillation waste product consisting of vapors of the isopropyl alcohol, water, and other contaminants with boiling points lower than isopropyl alcohol, pass through condenser 16 and into waste output line 72. Preferably, a differential pressure sensor 94 and a flow restrictor 96 are positioned on waste output line 72 to provide mass flow information for the column waste output line 72. Vapor waste flow through waste output line 72 is regulated by control valve 38, and waste vapors are directed to a waste product cooler 12, where they are cooled and condensed into liquid form. Cooling water from the water source line 86 flows into the waste product cooler 12 through an inlet water line 32. The flow of cooling water is controlled by the outlet water control valve 36. After passing through waste product cooler 12, the water is returned to the water return line 88 through the outlet water line 34. The resulting cool waste product passes through waste line 84.
to be stored and subsequently sent for off-site recycling. Preferably, a thermocouple 30 and a water concentration sensor 82 are positioned on waste line 84 to facilitate measurement of the temperature and water concentration of the waste product. The concentration sensor 82 may be an automatic concentration gauge based upon the dielectric constant of the isopropyl alcohol, or it may be an automatic Karl Fisher Titrator that determines the water content directly, or any other concentration monitoring means one skilled in the art may employ.

Figures 1 and 4 illustrate the details of the preferred embodiment of an apparatus suitable for carrying out the second distillation step of the present invention. In this step, the isopropyl alcohol is brought to an ultrapure level of zero particles per ml. larger than 2 microns, zero to 2 particles per ml. from 0.5 to 2 microns, zero to 30 particles per ml. from 0.1 to 0.5 microns, and unspecified number of particles less than 0.1 microns. The final product may contain any specific trace impurity, such as metals, cations, anions at levels from approximately 1 ppt to less than 1 ppb, and any other specific trace organic substances may be present at levels from 10 ppt to less than 10 ppm.

Referring now to Figure 4, second distillation sub-system 100 comprises a distillation column 164, a condenser 104, a reboiler 108, a reboiler waste product output line 160, a waste product cooler 106, and an end product cooler 102. Ultradry isopropyl alcohol from the first distillation sub-system 10, containing less than 100 ppm water and minor amounts of various high-boiling point contaminants, flows through a feed line 110 into the distillation column 164. Distillation column 164 uses high efficiency packing to provide from eleven to forty theoretical plates in the compact vertical space. Preferably, the distillation column 164 is packed such that it has twenty theoretical plates. In the twenty-plate embodiment, the isopropyl alcohol from the first distillation step is preferably introduced into distillation column 164 at point corresponding to
theoretical plate number fourteen out of twenty. The rectifying section 156 and stripping section 158 of distillation column 164 provide for the separation needed. Preferably, a pressure sensor 112 and a thermocouple 114 are positioned on the distillation column 164 near the isopropyl alcohol input point to facilitate the measurement of the pressure and temperature of the fluid within the distillation column 164.

The distillation column reboiler 108 provides a receptacle for fluids refluxing through the distillation column 164. The column reboiler 108 is heated by heat transfer from the reboiler steam coil 118. Steam coil 118 receives steam from a feed line 124, which in turn is connected to steam source line 90. Steam inlet control valve 126 controls the flow of steam through steam coil 118. Advantageously, adjustment of inlet control valve 126 allows for temperature regulation of the reboiler 108. Steam passing through steam coil 118 condenses, and the resulting steam condensate flows through the condensate return line 122 and into the steam return line 92. Preferably, a differential pressure sensor 116 is positioned on reboiler pressure line 188, where pressure line 188 connects the bottom of the distillation column 164 and the reboiler 108, to measure pressure differences. Also preferably, a thermocouple 120 is positioned within the reboiler 108 to facilitate temperature measurements of the fluid within the reboiler. A reboiler output line 160 is provided in the reboiler 108 to withdraw the waste product, consisting of isopropyl alcohol and various high boiling contaminants. Passing through the reboiler output line 160, the waste flows through purge flow metering pump 172, and into waste product cooler 106, where it is cooled and then channeled into the waste drain 694 for off-site recycling.

Distillation column condenser 104 is situated on top of the distillation column 164. Cooling water from the water source line 86 flows into the condensing coil 152 within the column condenser 104, causing the condensation of gaseous vapors rising up through the column, for proper
reflux back into column rectifying section 156. The cooling water exits condensing coil 152 through the condenser outlet water control valve 148 flowing to the water return line 88. Preferably, a thermocouple 146 and pressure sensor 144 are positioned with the condenser 104 to facilitate measurement of pressure and temperature.

Distillation end product, consisting of the vapor of ultrapure and ultradry isopropyl alcohol, passes through condenser 104 and into vapor product output line 162. Vapor product flow valve 142 regulates the flow of product into the vapor product output line 162. Preferably, a differential pressure sensor 174 and a flow restrictor 178 are positioned on vapor product output line 162 to provide mass flow information for the vapor product output line 162. Ultrapure and ultradry isopropyl alcohol vapor flows through the vapor product output line 162 to a vapor phase particle filter 170, which filters the isopropyl alcohol vapor to remove various contaminants distilled along with the isopropyl alcohol. After filtration in the vapor phase particle filter 170, the gaseous ultrapure and ultradry isopropyl alcohol enters the end product cooler 102, where it is cooled and condenses into liquid form. Cooling water flows into the end product cooler 102 through a inlet water line 138 from the water source line 86. The flow of cooling water is controlled by the outlet water control valve 136, and after passing through end product cooler 102, the water flows through outlet water line 140 into the water return line 88. The resulting cool ultrapure and ultradry isopropyl alcohol end product then passes through liquid product output line 134. Preferably, a thermocouple 154 and a water content monitor 168 are positioned on liquid product output line 134 to facilitate measurement of the temperature and water concentration of the ultrapure and ultradry isopropyl alcohol. The water content monitor 168 may be an automatic concentration gauge based the automatic Karl Fisher Titrator that determines the water content directly, or any other concentration monitoring means one skilled in the art may employ. For the very low
level of water content expected, a concentration monitor
based upon the dielectric constant of the isopropyl alcohol
is not preferred here.

In one preferred embodiment, illustrated in block
diagram form in Figure 1, the cooled ultrapure and ultradry
isopropyl alcohol is then sent to filtration and pumping
sub-system 300, through the end product output line 134,
for additional filtration. The final product is then
transported through a system isopropyl alcohol output line
310 to storage facilities for later use, or directly to the
semiconductor manufacturing process.

In another preferred embodiment, also illustrated in
Figure 1, a computer control system 200 monitors and
controls all aspects of the process through electrical and
signal connections to the various parts of the system.
Thus, for example, the computer controls the first
distillation sub-system 10 through data communication and
control instructions passing through data computer conduit
210. Similarly, the second distillation sub-system 100 is
controlled through computer conduit 220, the filtration and
pumping sub-system 300 through computer conduit 230, the
steam sub-system 400 through computer conduit 240, the
Control single line with cooling water sub-system 410
through computer conduit 250, and the pervaporation sub-
system 600 through computer conduit 260.

With respect to the computer controlled embodiment,
Figure 6 shows a set of control rules which may be executed
by the various hardware and software controllers present
within the system. The set of control rules comprise a
list of items to control, a list of sensors providing
control reference data, and a list of english description
of control rules.

Metering pump 650 is controlled by using data provided
by a pervaporation water content monitor 632 through rules
710 and 712. These rules provide for steady water
concentration of the partially dehydrated isopropyl alcohol
to output line 686. If the water content increases, then
the pump is slowed down to allow more time for the
pervaporation cell 620 to remove water.
Inlet steam control valve 628 is controlled by using data provided by a thermocouple 630 through the rules 714 and 716. These rules keep the temperature of the liquid in feed line 676 within the desired operating temperature range.

Water control valve 658 is controlled by using data provided by a thermocouple 636 through the rules 718 and 720. These rules keep the temperature of liquid in the waste water line 610 within the desired operating temperature range.

Metering pump 682 is controlled by using data provided by the level sensor 638 through rules 722 and 724. These rules keep the level of the liquid within the pervaporation waste water condenser 618 within the desired operating range.

Inlet steam control valve 28 is controlled by using data provided by thermocouple 74 through rules 728 and 730. These rules keep the temperature of the liquid in feed line 76 within the desired operating temperature range.

Outlet water control valve 36 is controlled by using data provided by a thermocouple 30 through rules 732 and 734. These rules keep the distillation waste 85 within the desired operating temperature range.

Control valve 38 is controlled by using data provided by concentration sensor 82 through the rules 736 and 738. These rules keep the concentration of distillation waste 85 slightly above the azeotropic concentration of isopropyl alcohol, thus ensuring most of the remaining water has been stripped from the liquid in feed line 76.

Reboiler inlet steam control valve 58 is controlled by using data provided by the differential pressure sensor 93 through rules 740 and 742. These rules keep the heat input to reboiler 18 at the proper setting to ensure proper boilup.

Reboiler inlet steam control valve 58 is also controlled by using data provided by pressure sensor 40 through rule 744. This rule prevents pressure runaway in column condenser 16.
Outlet water control valve 46 is controlled by using data provided by pressure sensor 40 through rules 746 and 748. These rules allow for proper regulation of column condenser 16.

Feed control valve 50 is controlled by using data provided by differential pressure sensor 66 through rules 750 and 752. These rules allow for proper flow of the ultradry isopropyl alcohol through output line 52 into feed line 110, and also prevent reboiler 18 from overfilling.

Inlet steam control valve 126 is controlled by using data provided by the differential pressure sensor 116 through rules 754 and 756. These rules keep the heat input to reboiler 108 at the proper setting to ensure proper boilup.

Inlet steam control valve 126 is also controlled by using data provided by differential pressure sensor 176 through rules 758 and 760. These rules keep the heat input to reboiler 108 at the proper setting to ensure proper boilup.

Outlet water control valve 148 is controlled by using data provided by the differential pressure sensor 95 through rules 762 and 764. These rules allow for proper regulation of column condenser 104.

Outlet water control valve 148 is also controlled by using data provided by pressure sensor 144 through rules 766 and 768. These rules allow for proper regulation of column condenser 104.

Control valve 142 is controlled by using data calculated from the mass flow data provided by feed flowmeter 80 minus the mass flow data from the purge flow metering pump 172 and compared to the mass flow data measured by the differential pressure sensor 174 through rules 770 and 772. These rules allow for proper material flow through the removal column 164.

Outlet water control valve 136 is controlled by using data calculated from thermocouple 154 through the rules 774 and 776. These rules keep the temperature of the liquid in product output line 134 within the desired operating temperature range.
Accordingly, the reader will see that we have invented a simpler and better method and process, allowing less complicated and lower cost equipment to be installed on-site at semiconductor manufacturing factories, for obtaining ultrapure and ultradry isopropyl alcohol.

Although the description of our invention contains many specificities, these should not be construed as limiting the scope of the invention but as merely providing illustrations of some of the presently preferred embodiments of this invention. For example, the high boiling contaminants could be removed first and the light boiling contaminants removed in the second distillation step of the process. The method and process as embodied in the preferred apparatus could be carried out in batch form of pervaporation and batch distillation steps, even allowing for the elimination of the second column, since the light contaminants could be removed first, then the same column could be used to purify the remaining dry isopropyl alcohol to ultrapure levels by taking the overhead condensate after the light impurities have been removed. Also, other preferred flow rates could allow a larger or smaller apparatus to be built employing the method and process of our invention.

Thus, the scope of the invention should be determined by the appended claims and their legal equivalents, rather than by the examples given.

Example 1.

Figure 5a shows the flow diagram of the preferred apparatus used in accordance with the process of the present invention for the conditions given below. Example 1 illustrates the use of the invention when the feed to the system contains less than a few thousand parts per million water content and therefore does not require a pervaporation stage to remove most of the water from the feed.

An isopropyl alcohol solution containing 1000 ppm n-propyl alcohol, approximately 500 ppm water, 10 ppm acetic acid, and 500 ppb metals, at 15 degrees Celsius is enters feed line 20 at a rate of approximately 36.29 kg/hr. The
solution is then conducted to the pre-heater column 14 where it is heated to 82 degrees Celsius with approximately 2.2 kW of heat supplied by the pre-heater column 14.

The heated isopropyl alcohol solution then enters distillation column 78 at a point corresponding to theoretical plate number fourteen out of twenty-five plates total. A light waste 85, consisting of approximately 88% isopropyl alcohol and any light boiling contaminants, is removed from the distillation column 78 at a rate of 0.4527 kg/hr with the use of 0.027 kW of cooling supplied to the waste product cooler 12 while 28.5 kW of cooling is supplied to the column condenser 16 to provide high level reflux of condensate back to the distillation column 78. 28.5 kW of heat is required in the column reboiler 18 to drive the first column.

35.83 kg/hr of now ultra-dry isopropyl alcohol is removed from the distillation column 78 and enters into distillation column 164 at a point corresponding to theoretical plate number fourteen out of twenty total theoretical plates. A heavy waste 133, consisting of approximately 75.9% isopropyl alcohol is sent to waste at a rate of 0.1425 kg/hr after being cooled by 0.024 kW of cooling supplied to the waste product cooler 106. Approximately 16.6 kW of heat is required in the column reboiler 108 to drive the second column.

The desired ultradry and now ultrapure isopropyl alcohol is delivered at a rate of approximately 35.696 kg/hr as the ultrapure and ultradry product 135, after it is cooled by 0.024 kW of cooling supplied to the end product cooler 102. The column condenser 104 supplies a high level of reflux for the removal column 164 when supplied with cooling of 16.6 kW. The overall efficiency of the two column distillation in this example is calculated to be 98.36% recovery of isopropyl alcohol.

Example 2.

Figure 5b shows the flow diagram of the preferred apparatus used in accordance with the process of the present invention for the conditions given below. Example 2 illustrates the use of the invention when the feed to the
system contains more than a few thousand parts per million water content and therefore requires a pervaporation stage to remove most of the water from the feed.

An isopropyl alcohol solution containing approximately 
1000 ppm n-propyl alcohol, 5% water, 10 ppm acetic acid, and 500 ppb metals, at approximately 15 degrees Celsius enters the pre-heater column 614 at a rate of 38.2 kg/hr where it is heated to 50 degrees Celsius with 1.1 kW of heat, and then enters the pervaporation cell 620. The pervaporation waste water condenser 618 removes approximately 1.9 kg/hr of a waste water from pervaporation condenser line 610. The waste water contains no more than 0.03 kg/hr of isopropyl alcohol.

Partially dehydrated isopropyl alcohol containing 1000 ppm n-propyl alcohol, approximately 500 ppm water, 10 ppm acetic acid, and 500 ppb metals, at 50 degrees Celsius exits the pervaporation sub-system 600 through output line 686 and enters the feed line 20 at a rate of approximately 36.29 kg/hr. The partially dehydrated isopropyl alcohol is then conducted to the pre-heater column 14 where it is heated to 82 degrees Celsius with approximately 1.1 kW of heat supplied to the pre-heater column 14.

The heated isopropyl alcohol enters the distillation column 78 at point corresponding to theoretical plate number fourteen out of twenty-five total. A light waste 85, consisting of approximately 88% isopropyl alcohol and any low boiling contaminants, is removed from the distillation column 78 at a rate of 0.4527 kg/hr with the use of 0.027 kW of cooling supplied to the waste product cooler 12. A total of 28.5 kW of cooling is supplied to the column condenser 16 to provide a high level reflux of condensate back to the distillation Column 78. A total of 28.5 kW of heat is required in the column reboiler 18 to drive the first column.

Ultradry isopropyl alcohol is removed from distillation column 78 at the rate of 35.83 kg/hr, and enters the distillation column 164 at a point corresponding to theoretical plate number fourteen of twenty total theoretical plates. A heavy waste 133, consisting of
approximately 75.9% of isopropyl alcohol is sent to waste at a rate of 0.1425 kg/hr after being cooled by 0.024 kW of cooling supplied to the waste product cooler 106. Approximately 16.6 kW of heat are required in the column reboiler 108 to drive the second column.

The desired ultradry and now ultrapure isopropyl alcohol end product is delivered at a rate of approximately 35.696 kg/hr after it is cooled by 0.024 kW of cooling supplied to the end product cooler 102. The column condenser 104 supplies a high level of reflux for the distillation column 164 when supplied with cooling of 16.6 kW. The overall efficiency of the two column distillation in this example yields a 98.36% recovery of isopropyl alcohol.

The present invention may be embodied in other specific forms without departing from its spirit or essential characteristics. The described embodiments are to be considered in all respects only as illustrative and not restrictive. The scope of the invention is, therefore, indicated by the appended claims rather than the foregoing description. All changes which come within the meaning and range of equivalency of the claims are to be embraced within their scope.
WHAT IS CLAIMED IS:

1. A method of dehydrating and purifying isopropyl alcohol from a solution containing isopropyl alcohol, water, and minor amounts of additional contaminants, resulting in an ultrapure and ultradry isopropyl alcohol, comprising the steps of:

   removing water from a solution containing isopropyl alcohol by pervaporation of the isopropyl alcohol solution through a water permeable membrane to produce a partially dehydrated isopropyl alcohol;

   removing substantially all of the remaining water and contaminants with boiling points less than isopropyl alcohol from the partially dehydrated isopropyl alcohol by distillation in an autonomous azeotropic self-stripping distillation column, and removing a partially purified and ultradry isopropyl alcohol from a reboiler at the bottom of the column;

   removing substantially all substances with boiling points greater than isopropyl alcohol from the ultradry and partially purified isopropyl alcohol by distillation through a low boiling overhead product distillation column, and taking an ultradry and ultrapure isopropyl alcohol end product as overhead condensate.

2. The method of Claim 1, wherein the autonomous azeotropic self-stripping distillation column comprises eleven to forty theoretical distillation plates.

3. The method of Claim 1, wherein the low boiling overhead product distillation column comprises eleven to forty theoretical distillation plates.

4. The method of Claim 1, further comprising filtering the ultrapure and ultradry isopropyl alcohol end product to remove submicron sized particulate impurities therefrom.

5. The method of Claim 1, wherein the ultrapure and ultradry isopropyl alcohol end product contains less than about 100 parts per million water, zero particles per milliliter of a size greater than two microns, less than two particles per milliliter of a size greater than 0.5
microns but less than 2.0 microns, less than about thirty particles per milliliter of a size less than 0.5 micron but greater than 0.1 micron, and having less than about 1 part per billion of any specific trace inorganic impurity, and less than about ten parts per million of any specific trace organic impurity.

6. A method of dehydrating and purifying isopropyl alcohol from a solution containing isopropyl alcohol, and minor amounts of additional organic and inorganic contaminants, and less than about 2000 parts per million water, comprising the steps of:

   removing substantially all of the water and contaminants with boiling points less than isopropyl alcohol from the solution containing less than about 2000 parts per million water by distilling the solution in an autonomous azeotropic self-stripping distillation column, and removing a partially purified and ultradry isopropyl alcohol from a reboiler at the column bottom;

   removing substantially all substances with boiling points greater than isopropyl alcohol from the ultradry and partially purified isopropyl alcohol by distilling the ultradry and partially purified isopropyl alcohol through a low boiling overhead product distillation column, taking an ultradry and ultrapure isopropyl alcohol end product as overhead condensate.

7. The method of Claim 6, wherein the autonomous azeotropic self-stripping distillation column comprises eleven to forty theoretical distillation plates.

8. The method of Claim 6, wherein the low boiling overhead product distillation column comprises eleven to forty theoretical distillation plates.

9. The method of Claim 6, further comprising filtering the ultrapure and ultradry isopropyl alcohol end product to remove particulate impurities therefrom.

10. The method of Claim 6, wherein the ultrapure and ultradry isopropyl alcohol end product contains less than
about 100 parts per million water, zero particles per
milliliter of a size greater than two microns, less than
two particles per milliliter of a size greater than 0.5
microns but less than 2.0 microns, less than about thirty
particles per milliliter of a size less than 0.5 micron but
greater than 0.1 micron, and having less than about 1 part
per billion of any specific trace inorganic impurity, and
less than about ten parts per million of any specific trace
organic impurity.

11. An apparatus for dehydrating and purifying
isopropyl alcohol from a solution containing isopropyl
alcohol, water, and various other contaminants, the
apparatus comprising:
   a isopropyl alcohol feed line;
   means for pervaporation of isopropyl alcohol, the
   pervaporation means connected to the isopropyl alcohol
   feed line as an input line and having an output line;
   a first distillation system, the first
   distillation system comprising a pre-heating column
   connected to the pervaporation means output line, an
   autonomous azeotropic self-stripping distillation
   column connected to the pre-heating column, a
   condenser connected to the top of the autonomous
   azeotropic self-stripping distillation column, a
   reboiler connected to the bottom of the autonomous
   azeotropic self-stripping distillation column, and a
   reboiler output line connected to the reboiler for
   removing ultradry isopropyl alcohol;
   a second distillation system comprising a low
   boiling overhead product distillation column connected
   to the reboiler output line, a reboiler connected to
   the bottom of the low boiling overhead product
   distillation column, an end product condenser
   connected to the top of the low boiling overhead
   product distillation column, a product cooling column
   connected to the end product condenser, and an end
   product output line connected to the end product
   condenser.

12. The apparatus of Claim 11, wherein the autonomous
azeotropic self-stripping distillation column comprises
11. The apparatus of Claim 11, wherein the low
eleven to forty theoretical plates.
boiling overhead product distillation column comprises
14. The apparatus of Claim 11, further comprising a
eleven to forty theoretical plates.
filter connected to the end product output line, the filter
being capable of removing submicron sized particulate
impurities from isopropyl alcohol passing therethrough.
15. An apparatus for dehydrating and purifying
isopropyl alcohol from a solution containing isopropyl
alcohol, minor amounts of additional organic and inorganic
contaminants, and less than about 2000 parts per million
water, the apparatus comprising:

- a isopropyl alcohol feed line;
- a first distillation system, the first
distillation system comprising a pre-heating column
connected to the isopropyl alcohol feed line, an
autonomous azeotropic self-stripping distillation
column connected to the pre-heating column, a
condenser situated at the top of the autonomous
azeotropic self-stripping distillation column, a
reboiler situated at the bottom of the autonomous
azeotropic self-stripping distillation column, and a
reboiler output line connected to the reboiler for
removing ultradry isopropyl alcohol;
- a second distillation system comprising a low
boiling overhead product distillation column connected
to the reboiler output line, a reboiler situated at
the bottom of the low boiling overhead product
distillation column, an end product condenser situated
at the top of the low boiling overhead product
distillation column, a product cooling column
connected to the end product condenser, and an end
product output line connected to the end product
condenser.
16. The apparatus of Claim 15, wherein the autonomous
azeotropic self-stripping distillation column comprises
eleven to forty theoretical plates.
17. The apparatus of Claim 15, wherein the low boiling overhead product distillation column comprises eleven to forty theoretical plates.

18. The apparatus of Claim 15, further comprising filtration means connected to the end product output line, the filtration means being capable of removing submicron sized particulate impurities from isopropyl alcohol passing therethrough.
AMENDED CLAIMS

[received by the International Bureau on 22 November 1996 (22.11.96); original claims 1-13 and 15-17 amended; remaining claims unchanged (5 pages)]

1. A method of dehydrating and purifying isopropyl alcohol from a solution containing isopropyl alcohol, water, and impurities selected from the group consisting of metals, anions, cations, organic substances with a lower boiling point than isopropyl alcohol, organic substances with a higher boiling point than isopropyl alcohol and particulate matter, resulting in an ultrapure and ultradry isopropyl alcohol, said method comprising the steps of:

   10 removing water from a solution containing isopropyl alcohol by pervaporation of the isopropyl alcohol solution through a water permeable membrane to produce a partially dehydrated isopropyl alcohol;

   15 removing substantially all of the remaining water and impurities with boiling points less than isopropyl alcohol from the partially dehydrated isopropyl alcohol by a first distillation in a stripping distillation column, and removing a partially purified and ultradry isopropyl alcohol from a reboiler at the bottom of the column;

   20 removing substantially all substances with boiling points greater than isopropyl alcohol from the ultradry and partially purified isopropyl alcohol by a second distillation through a low boiling overhead product distillation column, and taking an ultradry and ultrapure isopropyl alcohol end product as overhead product.

2. The method of Claim 1, wherein the first distillation column comprises eleven to forty theoretical distillation plates.

3. The method of Claim 1, wherein the second distillation column comprises eleven to forty theoretical distillation plates.

4. The method of Claim 1, further comprising filtering the ultrapure and ultradry isopropyl alcohol end product to remove submicron sizes of said particulate matter.

5. The method of Claim 1, wherein the ultrapure and ultradry isopropyl alcohol end product contains less than
about 100 parts per million water, zero particles per milliliter of a size greater than two microns, less than two particles per milliliter of a size greater than 0.5 microns, less than about thirty particles per milliliter of a size less than 0.5 micron but greater than 0.1 micron, and having less than about 1 part per billion of any specific anion or cation, and less than about ten parts per million of any specific organic substance with a boiling point lower or higher than isopropyl alcohol.

6. A method of dehydrating and purifying isopropyl alcohol from a solution containing isopropyl alcohol, less than about 2000 parts per million water, impurities selected from the group consisting of metals, anions, cations, organic substances with a lower boiling point than isopropyl alcohol, organic substances with a higher boiling point than isopropyl alcohol and particulate matter, resulting in an ultrapure and ultradry isopropyl alcohol, comprising the steps of:

   removing substantially all of the water and contaminants with boiling points less than isopropyl alcohol from the solution containing less than about 2000 parts per million water by a first distillation in a stripping distillation column, and removing a partially purified and ultradry isopropyl alcohol from a reboiler at the column bottom;

   removing substantially all impurities with boiling points greater than isopropyl alcohol from the ultradry and partially purified isopropyl alcohol by a second distillation through a low boiling overhead product distillation column, and taking an ultradry and ultrapure isopropyl alcohol end product as overhead product.

7. The method of Claim 6, wherein the first distillation column comprises eleven to forty theoretical distillation plates.

8. The method of Claim 6, wherein the second distillation column comprises eleven to forty theoretical distillation plates.

9. The method of Claim 6, further comprising filtering
the ultrapure and ultradry isopropyl alcohol end product to
remove submicron sizes of said particulate matter.

10. The method of Claim 6, wherein said ultrapure and
ultradry isopropyl alcohol end product contains less than
about 100 parts per million water, zero particles per
milliliter of a size greater than two microns, less than two
particles per milliliter of a size greater than 0.5 microns,
less than about thirty particles per milliliter of a size less
than 0.5 micron but greater than 0.1 micron, and having less
than about 1 part per billion of any specific anion or cation,
and less than about ten parts per million of any specific
organic substance with a lower or higher boiling point than
isopropyl alcohol.

11. An apparatus for dehydrating and purifying isopropyl
alcohol from a solution containing isopropyl alcohol, water,
and impurities selected from the group consisting of metals,
anions, cations, organic substances with a lower boiling point
than isopropyl alcohol, organic substances with a higher
boiling point than isopropyl alcohol and particulate matter,
resulting in an ultrapure and ultradry isopropyl alcohol, the
apparatus comprising:

   a isopropyl alcohol feed line;
   means for pervaporation of isopropyl alcohol, the
   pervaporation means connected to the isopropyl alcohol
   feed line as an input line and having an output line;
   a first distillation system, the first distillation
   system comprising a pre-heating apparatus connected to
   the pervaporation means output line, a first distillation
   column connected to the pre-heating apparatus, a
   condenser connected to the top of the first distillation
   column, a reboiler connected to the bottom of the first
   distillation column, and a reboiler output line connected
to the reboiler for removing ultradry isopropyl alcohol;
   a second distillation system comprising a low
   boiling overhead product distillation column connected
to the reboiler output line, a reboiler connected to the bottom
   of the low boiling overhead product distillation
column, an end product condenser connected to the top of the low boiling overhead product distillation column, a product cooling column connected to the end product condenser, and an end product output line connected to the end product condenser.

12. The apparatus of Claim 11, wherein the first distillation column comprises eleven to forty theoretical plates.

13. The apparatus of Claim 11, wherein the second distillation column comprises eleven to forty theoretical plates.

14. The apparatus of Claim 11, further comprising a filter connected to said end product output line, the filter being capable of removing submicron sized particulate impurities from isopropyl alcohol passing therethrough.

15. An apparatus for dehydrating and purifying isopropyl alcohol from a solution containing isopropyl alcohol, less than about 2000 parts per million water, impurities selected from the group consisting of metals, anions, cations, organic substances with a lower boiling point than isopropyl alcohol, organic substances with a higher boiling point than isopropyl alcohol and particulate matter, resulting in an ultrapure and ultradry isopropyl alcohol and the apparatus comprising:

   a isopropyl alcohol feed line;

   a first distillation system, the first distillation system comprising a pre-heating apparatus connected to the isopropyl alcohol feed line, a first distillation column connected to the pre-heating apparatus, a condenser connected to the top of the first distillation column, a reboiler connected to the bottom of the first distillation column, and a reboiler output line connected to the reboiler for removing ultradry isopropyl alcohol;

   a second distillation system comprising a low boiling overhead product distillation column connected to the reboiler output line, a reboiler connected to the bottom of the low boiling overhead product distillation column, an end product condenser connected to the top of
the low boiling overhead product distillation column, a product cooling column connected to the end product condenser, and an end product output line connected to the end product condenser.

16. The apparatus of Claim 15, wherein the first distillation column comprises eleven to forty theoretical distillation plates.

17. The apparatus of Claim 15, wherein the second distillation column comprises eleven to forty theoretical plates.

18. The apparatus of Claim 15, further comprising filtration means connected to said end product output line, the filtration means being capable of removing submicron sized particulate impurities from isopropyl alcohol passing therethrough.
Statement under PCT Article 19(1)

The claims as amended recite an apparatus and method for removal of impurities from isopropyl alcohol, including organic impurities, by pervaporation followed by two distillation steps. Kagiyama et al. do not teach the removal of organic impurities and only use a single distillation column for production of ultrapure isopropyl alcohol. Further, Kagiyama et al. do not teach production of an ultradry isopropyl alcohol preparation because water is not stripped off during the distillation step as shown in Examples 1 and 3 (99.2% isopropyl alcohol after pervaporation versus 99.1% after distillation and 99.7% isopropyl alcohol after both pervaporation and after distillation, respectively).

Pasternak et al. disclose removal of water from concentrated solutions by pervaporation. Example XXVII, Col. 9, yields 1.1% of the isopropyl alcohol in the permeate and a separation factor of 809 which translates to greater than 1% water in the remaining isopropyl alcohol. Accordingly, Pasternak et al. do not teach production of ultrapure isopropyl alcohol, but are only concerned with recovering a product containing decreased amounts of water (See Col. 3, lines 28-31). Further, the process of Pasternak et al. may involve distillation using only a single distillation column.
FIG. 1

SUBSTITUTE SHEET (RULE 26)
<table>
<thead>
<tr>
<th>Item to Control</th>
<th>Sensor number</th>
<th>Rule number</th>
<th>Rule Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>650</td>
<td>632</td>
<td>710</td>
<td>If water content is &gt; 500 ppm slow pump rate</td>
</tr>
<tr>
<td>650</td>
<td>632</td>
<td>712</td>
<td>If water content &lt; 300 ppm increase pump rate</td>
</tr>
<tr>
<td>628</td>
<td>630</td>
<td>714</td>
<td>If temperature &gt; T-pervep + 10% then decrease flow in valve</td>
</tr>
<tr>
<td>628</td>
<td>630</td>
<td>716</td>
<td>If temperature &lt; T-pervep - 10% then increase flow in valve</td>
</tr>
<tr>
<td>658</td>
<td>636</td>
<td>718</td>
<td>If temperature &gt; T-Waste + 10% then increase flow in valve</td>
</tr>
<tr>
<td>658</td>
<td>636</td>
<td>720</td>
<td>If temperature &lt; T-Waste - 10% then decrease flow in valve</td>
</tr>
<tr>
<td>658</td>
<td>638</td>
<td>722</td>
<td>If level &gt; L-Waste + 10% then increase pump rate</td>
</tr>
<tr>
<td>658</td>
<td>638</td>
<td>724</td>
<td>If level &lt; L-Waste - 10% then decrease pump rate</td>
</tr>
<tr>
<td>28</td>
<td>74</td>
<td>728</td>
<td>If temperature &gt; T-preheat + 10% then decrease flow in valve</td>
</tr>
<tr>
<td>28</td>
<td>74</td>
<td>730</td>
<td>If temperature &lt; T-preheat - 10% then increase flow in valve</td>
</tr>
<tr>
<td>36</td>
<td>30</td>
<td>732</td>
<td>If temperature &gt; T-cold-waste + 10% then increase flow in valve</td>
</tr>
<tr>
<td>36</td>
<td>30</td>
<td>734</td>
<td>If temperature &lt; T-cold-waste - 10% then decrease flow in valve</td>
</tr>
<tr>
<td>36</td>
<td>82</td>
<td>736</td>
<td>If concentration &lt; Azeotrope + 2% then decrease flow in valve</td>
</tr>
<tr>
<td>36</td>
<td>82</td>
<td>738</td>
<td>If concentration &gt; Azeotrope + 1% then increase flow in valve</td>
</tr>
<tr>
<td>58</td>
<td>93</td>
<td>740</td>
<td>If pressure difference &gt; p-light-column + 2% then decrease flow in valve</td>
</tr>
<tr>
<td>58</td>
<td>93</td>
<td>742</td>
<td>If pressure difference &lt; p-light-column - 2% then increase flow in valve</td>
</tr>
<tr>
<td>58</td>
<td>40</td>
<td>744</td>
<td>If absolute pressure &gt; 5 psig then decrease flow in valve</td>
</tr>
<tr>
<td>46</td>
<td>40</td>
<td>746</td>
<td>If absolute pressure &gt; 2 psig then increase flow in valve</td>
</tr>
<tr>
<td>46</td>
<td>40</td>
<td>748</td>
<td>If absolute pressure &lt; 1 psig then decrease flow in valve</td>
</tr>
<tr>
<td>50</td>
<td>66</td>
<td>750</td>
<td>If differential pressure &gt; p-light-boil + 5% then increase flow in valve</td>
</tr>
<tr>
<td>50</td>
<td>66</td>
<td>752</td>
<td>If differential pressure &lt; p-light-boil - 5% then decrease flow in valve</td>
</tr>
<tr>
<td>126</td>
<td>116</td>
<td>754</td>
<td>If differential pressure &gt; p-heavy-boil + 5% then increase flow in valve</td>
</tr>
<tr>
<td>126</td>
<td>116</td>
<td>756</td>
<td>If differential pressure &lt; p-heavy-boil - 5% then decrease flow in valve</td>
</tr>
<tr>
<td>126</td>
<td>176</td>
<td>758</td>
<td>If differential pressure &gt; p-heavy-column + 5% then decrease flow in valve</td>
</tr>
<tr>
<td>126</td>
<td>176</td>
<td>760</td>
<td>If differential pressure &lt; p-heavy-column - 5% then increase flow in valve</td>
</tr>
<tr>
<td>148</td>
<td>95</td>
<td>762</td>
<td>If differential pressure &gt; p-light-heavy + 5% then decrease flow in valve</td>
</tr>
<tr>
<td>148</td>
<td>95</td>
<td>764</td>
<td>If differential pressure &lt; p-light-heavy - 5% then increase flow in valve</td>
</tr>
<tr>
<td>148</td>
<td>144</td>
<td>766</td>
<td>If absolute pressure &gt; 1 psig then increase flow in valve</td>
</tr>
<tr>
<td>148</td>
<td>144</td>
<td>768</td>
<td>If absolute pressure &lt; 0.5 psig then decrease flow in valve</td>
</tr>
<tr>
<td>142</td>
<td>80 - 172</td>
<td>770</td>
<td>If flow rate of 80 minus rate of 172 &gt; flow rate measured by 174 + 1% then increase flow in valve</td>
</tr>
<tr>
<td>142</td>
<td>80 - 172</td>
<td>772</td>
<td>If flow rate of 80 minus rate of 172 &lt; flow rate measured by 174 - 1% then decrease flow in valve</td>
</tr>
<tr>
<td>136</td>
<td>154</td>
<td>774</td>
<td>If temperature &gt; T-product +5% then increase flow in valve</td>
</tr>
<tr>
<td>136</td>
<td>154</td>
<td>776</td>
<td>If temperature &lt; T-product -5% then decrease flow in valve</td>
</tr>
</tbody>
</table>

**FIG. 6**

SUBSTITUTE SHEET (RULE 26)
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) :B01D 3/00; C07C 29/80; 31/18
US Cl :Please See Extra Sheet.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 159/Dig. 27; 202/154, 155, 172, 173, 176, 202; 203/18, 71, Dig. 16; 210/295, 640; 568/889, 916

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category*</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y</td>
<td>US,A, 5,182,022 (PASTERNAK ET AL) 26 January 1993 see column 2, lines 56-64 and column 3, lines 12-27</td>
<td>1, 6, 11, 15</td>
</tr>
<tr>
<td>Y</td>
<td>US,A, 4,788,043 (KAGIYAMA ET AL) 29 November 1988, See the Abstract and column 9, lines 60-63</td>
<td>1, 4, 6, 9, 11, 14-15, 18</td>
</tr>
<tr>
<td>A</td>
<td>GB,A, 856,371 (BINNING ET AL) 14 December 1960, see entire document</td>
<td>1-18</td>
</tr>
</tbody>
</table>

* Special categories of cited documents:
  *T* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
  *T* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
  *X* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
  *Y* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
  *Z* document member of the same patent family

Date of the actual completion of the international search
29 AUGUST 1995

Date of mailing of the international search report
28 SEP 1995

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Telephone No. (703) 308-3844

Form PCT/ISA/210 (second sheet)(July 1992)
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER:
US CL :

159/Dig. 27; 202/155, 173, 176, 202; 203/18, 17, Dig. 16; 210/295, 640; 568/889, 916

Form PCT/ISA/210 (extra sheet)(July 1992)