

METHANOL CARBONYLATION SYSTEM HAVING ABSORBER WITH
MULTIPLE SOLVENT OPTIONS

5 Claim for Priority

 This non-provisional application claims the benefit of the filing date of United States Provisional Patent Application Serial No. 61/125,791, of the same title, filed April 29, 2009. The priority of United States Provisional Patent Application Serial No. 61/125,791 is hereby claimed and the disclosure thereof is
10 incorporated into this application by reference.

Technical Field

 The present invention relates to acetic acid production and, in particular, to a methanol carbonylation system with a light ends absorber adapted to use
15 different scrubber solvents and return the used solvent to the carbonylation system.

Background of the Invention

 Acetic acid production by way of methanol carbonylation is known in the
20 art. Generally speaking, a methanol carbonylation production line includes a reaction section, a purification section, light ends recovery and a catalyst reservoir system. In the reaction section, methanol and carbon monoxide are contacted with rhodium or iridium catalyst in a homogenous stirred liquid phase reaction medium in a reactor to produce acetic acid. Methanol is pumped to the reactor from a
25 methanol surge tank. The process is highly efficient, having a conversion of methanol to acetic acid of typically greater than 99 percent. The reaction section also includes a flash vessel coupled to the reactor which flashes a draw stream in order to remove crude product from the reaction section. The crude product is fed to a purification section which includes generally a light ends or stripper column,
30 a drying column, auxiliary purification and optionally a finishing column. In the process, various vent streams containing light ends, notably methyl iodide, carbon monoxide and methyl acetate are generated and fed to the light ends recovery

section. These vent streams are scrubbed with a solvent to remove the light ends which are returned to the system or discarded.

In a traditional, *Monsanto* methanol carbonylation plant, a high pressure
5 and low pressure absorber are included wherein acetic acid is used as the scrubber solvent. The acetic acid solvent must subsequently be stripped of light ends, usually in another purification column so that the acid is not wasted. Such columns are expensive because they must be made of a highly corrosion resistant material such as zirconium alloys and so forth. Moreover, stripping light ends
10 from the acid requires steam and contributes to operating expense.

Nevertheless, using acetic acid as a scrubber solvent is widespread in the carbonylation art generally. *See* for example, United States Patent No. 5,502,243 to *Waller et al.*, entitled "Hydrocarbonylation of Dimethyl Ether". *Note* the
15 disclosure at **Figure 3**, and the discussion at Cols. 8 and 9 concerning operation of an absorber **321**. A cool acetic stream **323** passes downwardly through this absorber and absorbs any residual co-products and volatile catalyst components from the vent gas.

20 *So also*, there is disclosed in United States Patent No. 4,241,219 to *Wan*, entitled "Treatment of Carbonylation Effluent", a method of recovering volatile components by contact with a scrubbing solvent recovered from the reaction mixture in the same production line. *See* Col. 2, lines 15-30 wherein it is noted that acetic anhydride, ethylidene diacetate, acetic acid, or mixtures of them can be
25 used as a vent gas scrubber solvent.

Methanol has been suggested for use as a scrubber solvent in connection with a methanol carbonylation process. In this regard, *see* United States Patent No. 5,416,237 to *Aubigne et al.*, entitled "Process for the Production of Acetic
30 Acid". It is noted in the '237 patent that noncondensables from a flash tank vapor overhead may be scrubbed countercurrently with chilled methanol. The methanol

scrubber solvent residual stream is added to pure methanol and then used as feed to the reactor. *See* Col. 9, lines 30-42. If the reactor is not consuming the residual stream, it must be stored separately or purified again contributing to capital expense and operating costs.

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Chinese Patent Application Publication No. 200410016120.7 discloses a method for recovering light components in vent gas from production of acetic acid/acetic anhydride by way of scrubbing with methanol and acetic acid. The system disclosed in the Publication No. 200410016120.7 discloses a two stage
10 absorption arrangement wherein vent gas is treated sequentially with two different absorbents in a two stage system. *Note* particularly **Figure 2**. Another system is seen in an industrial publication entitled "Process of 200ktpa Methanol Low Press Oxo Synthesis AA" (SWRDICI 2006) (China). In this research publication there is shown a vent gas treatment system including a high pressure absorber as well as
15 a low pressure absorber. Both absorbers of this system are operated utilizing methanol as a scrub fluid.

While there have been advances in the art, known methods of scrubbing vent gases in methanol carbonylation systems typically involve multiple absorber
20 towers which are expensive to fabricate and operate. In accordance with the invention, there is provided an improved methanol carbonylation system with an absorber capable of using different solvents. The inventive system reduces both capital requirements and operating costs as compared with conventional systems.

25 Summary of the Invention

There is provided a carbonylation system for making acetic acid having an absorber tower with multiple scrubber solvent options for treating vent gas. The absorber recovers methyl iodide and other volatiles such as methyl acetate vapor from the vent gas with scrubber solvent, the tower being coupled to first and
30 second scrubber solvent sources which are capable of supplying different scrubber solvents. Typically, methanol is used as a scrubber solvent in a steady state mode

of operation, while acetic acid may be used during start-up or transient operation of the unit. A switching system alternatively provides either methanol or acetic acid to the tower and returns the solvent and recovered volatiles to the carbonylation system for further reaction. During changeover of scrubber
5 solvents, the recovered material may be added to the catalyst reservoir system if so desired.

The use of a one column system light ends absorber in accordance with the invention allows operation without the need for a dedicated light ends stripper for
10 recovering methyl iodide from the system offgas vent stream. A salient benefit is capital reduction for new acetic acid carbonylation projects (eliminate stripper column system, reboiler, subcooler, and associated instrumentation and piping) by being able to use one absorber system on two different scrub mediums to accommodate all modes of operation including: startup, normal operation, upset
15 operation and shutdown. Another benefit is energy reduction for normal operation via steam savings realized by elimination of the need for an auxiliary stripper column for scrubber solvent. Still another benefit is better scrub of light ends component methyl iodide by using sub-chilled 5-15 °C methanol for a normal operation mode.

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Further details and advantages will be apparent from the discussion which follows.

Brief Description of the Drawings

25 The invention is described in detail below with reference to the appended drawings, wherein like numerals designate similar parts. In the drawings:

Figure 1 is a schematic diagram illustrating a carbonylation system for making acetic acid and

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Figure 2 is a schematic diagram illustrating a vent gas absorber and switching system used in connection with the system of **Figure 1**.

Detailed Description of the Invention

5 The invention is described in detail below with reference to numerous embodiments for purposes of exemplification and illustration only. Modifications to particular embodiments within the spirit and scope of the present invention, set forth in the appended claims, will be readily apparent to those of skill in the art.

10 Unless more specifically defined below, terminology as used herein is given its ordinary meaning. % and like terms refer to weight percent, unless otherwise indicated.

 “Consisting essentially of” and like terminology refers to a composition
15 consisting of 90% by weight or more of the specified component. Thus a scrubber solvent stream consists essentially of methanol if it is at least 90% methanol.

 “Low pressure” and like terminology refers to pressures much lower than the pressure maintained in a carbonylation reactor of the class discussed herein.
20 Low pressure thus refers to gauge pressures generally less than 10 bar, suitably less than 5 bar, typically less than about 3 bar and in some cases less than 1 bar.

 “Volatile” components are those compounds in vapor phase and/or having a boiling point below or equal to that of methyl acetate including methyl iodide.
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 As used herein the “purified process stream” includes the process stream fed forward from the light ends column, and any subsequent purifications of the light ends process stream.

30 A Group VIII catalyst metal used in connection with the present invention may be a rhodium and/or iridium catalyst. The rhodium metal catalyst may be

added in any suitable form such that rhodium is in the catalyst solution as an equilibrium mixture including $[\text{Rh}(\text{CO})_2\text{I}_2]^-$ anion as is well known in the art. When rhodium solution is in the carbon monoxide-rich environment of the reactor, solubility of the rhodium is generally maintained because

5 rhodium/carbonyl iodide anionic species are generally soluble in water and acetic acid. However, when transferred to carbon monoxide depleted environments as typically exist in the flasher, light ends column and so forth, the equilibrium rhodium/catalyst composition changes since less carbon monoxide is available. Rhodium precipitates as RhI_3 , for example; details as to the form of entrained

10 rhodium downstream of the reactor is not well understood. Iodide salts help alleviate precipitation in the flasher under so-called "low water" conditions as will be appreciated by one of skill in the art.

Iodide salts maintained in the reaction mixtures of the processes described

15 herein may be in the form of a soluble salt of an alkali metal or alkaline earth metal or a quaternary ammonium or phosphonium salt. In certain embodiments, the catalyst co-promoter is lithium iodide, lithium acetate, or mixtures thereof. The iodide salt may be added as a mixture of salts such as a mixture of lithium iodide and sodium iodide and/or potassium iodide. Alternatively, the iodide salt

20 may be generated in-situ since under the operating conditions of the reaction system, a wide range of non-iodide salt precursors such as alkali metal acetates will react with methyl iodide to generate the corresponding co-promoter iodide salt stabilizer. Suitable salts can be generated *in situ* from non-ionic precursors, such as a phosphine oxide or any suitable organic ligand or ligands if so desired.

25 Phosphine oxides and suitable organic ligands generally undergo quaternization in the presence of methyl iodide at elevated temperatures to yield salts which maintain iodide anion concentration. For additional detail regarding iodide salt generation, *see* United States Patent Nos. 5,001,259 to *Smith et al.*; 5,026,908 to *Smith et al.*; and 5,144,068, also to *Smith et al.*, the disclosures of which are

30 hereby incorporated by reference.

An iridium catalyst in the liquid carbonylation reaction composition may comprise any iridium-containing compound which is soluble in the liquid reaction composition. The iridium catalyst may be added to the liquid reaction composition for the carbonylation reaction in any suitable form which dissolves in the liquid reaction composition or is convertible to a soluble form. Examples of suitable iridium-containing compounds which may be added to the liquid reaction composition include: IrCl_3 , IrI_3 , IrBr_3 , $[\text{Ir}(\text{CO})_2\text{I}]_2$, $[\text{Ir}(\text{CO})_2\text{Cl}]_2$, $[\text{Ir}(\text{CO})_2\text{Br}]_2$, $[\text{Ir}(\text{CO})_2\text{I}_2]^- \text{H}^+$, $[\text{Ir}(\text{CO})_2\text{Br}_2]^- \text{H}^+$, $[\text{Ir}(\text{CO})_2\text{I}_4]^- \text{H}^+$, $[\text{Ir}(\text{CH}_3)_3(\text{CO}_2)]^- \text{H}^+$, $\text{Ir}_4(\text{CO})_{12}$, $\text{IrCl}_3 \cdot 3\text{H}_2\text{O}$, $\text{IrBr}_3 \cdot 3\text{H}_2\text{O}$, $\text{Ir}_4(\text{CO})_{12}$, iridium metal, Ir_2O_3 , $\text{Ir}(\text{acac})(\text{CO})_2$, $\text{Ir}(\text{acac})_3$, iridium acetate, $[\text{Ir}_3\text{O}(\text{OAc})_6(\text{H}_2\text{O})_3][\text{OAc}]$, and hexachloroiridic acid $[\text{H}_2\text{IrCl}_6]$. Chloride-free complexes of iridium such as acetates, oxalates and acetoacetates are usually employed as starting materials. The iridium catalyst concentration in the liquid reaction composition may be in the range of 100 to 6000 ppm. The carbonylation of methanol utilizing iridium catalyst is well known and is generally described in the following United States Patents: 5,942,460; 5,932,764; 5,883,295; 5,877,348; 5,877,347 and 5,696,284, the disclosures of which are hereby incorporated by reference into this application as if set forth in their entirety.

A supported Group VIII catalyst may employed if so desired. One preferred system includes an insoluble polymer having pendent pyrrolidone groups which support a rhodium species. One suitable catalyst is a poly-vinylpyrrolidone which has been crosslinked and rhodium loaded. Cross-linking can be achieved using a caustic catalyst as disclosed in United States Patent No. 2,938,017 or by using a cross-linking agent such as disclosed in German 2,059,484. These references are herein incorporated by reference. This catalyst is prepared by reacting the polymer support with an alkyl halide and a rhodium compound. Both reactions are readily accomplished by standard procedures and using known components for such reactions. For example, it is preferred to simply add an amount of the insoluble polymer such as in powder or resin bead form to what otherwise constitutes as the homogeneous medium for the methanol

carbonylation reaction. Such carbonylation reaction medium includes methanol and/or methyl acetate, acetic acid and a small amount of water in a pressure vessel along with a rhodium compound and an iodide promoter as described herein. Further details appear in United States Patent No. 5,466,874, the disclosure of which is incorporated herein by reference in its entirety.

Another system includes an insoluble, pyridine ring-containing polymer, and a Group VIII metal supported thereon and is known *per se*. The term "pyridine ring-containing polymer" used herein is intended to refer to a polymer containing substituted or non-substituted pyridine rings or substituted or non-substituted, pyridine-containing polycondensed rings such as quinoline rings. The substituents include those inert to the methanol carbonylation such as an alkyl group and alkoxy group. Typical examples of the insoluble, pyridine ring-containing polymers include those obtained by reaction of vinylpyridine with a divinyl monomer or by reaction of vinylpyridine with a divinyl monomer-containing vinyl monomer, such as copolymers of 4-vinylpyridine and divinylbenzene, copolymers of 2-vinylpyridine and di-vinylbenzene, copolymers of styrene, vinylbenzene and divinylbenzene, copolymers of vinylmethylpyridine and divinylbenzene and copolymers of vinylpyridine, methyl acrylate and ethyl diacrylate. Further details appear in United States Patent No. 5,334,755, the disclosure of which is incorporated herein by reference in its entirety.

Methyl iodide is used as the promoter. Preferably, the concentration of methyl in the liquid reaction composition is in the range 1 to 50% by weight, preferably 2 to 30% by weight.

The promoter may be combined with a salt stabilizer/co-promoter compound, which may include salts of a metal of Group IA or Group IIA, or a quaternary ammonium or phosphonium salt. Particularly preferred are iodide or acetate salts, *e.g.*, lithium iodide or lithium acetate.

Other promoters and co-promoters may be used as part of the catalytic system of the present invention as described in European Patent Publication EP 0 849 248, the disclosure of which is hereby incorporated by reference. Suitable promoters are selected from ruthenium, osmium, tungsten, rhenium, zinc, 5 cadmium, indium, gallium, mercury, nickel, platinum, vanadium, titanium, copper, aluminum, tin, antimony, and are more preferably selected from ruthenium and osmium. Specific co-promoters are described in United States Patent No. 6,627,770, the entirety of which is incorporated herein by reference.

10 A promoter may be present in an effective amount up to the limit of its solubility in the liquid reaction composition and/or any liquid process streams recycled to the carbonylation reactor from the acetic acid recovery stage. When used, the promoter is suitably present in the liquid reaction composition at a molar ratio of promoter to metal catalyst of [0.5 to 15]:1, preferably [2 to 10]:1, more 15 preferably [2 to 7.5]:1. A suitable promoter concentration is 400 to 5000 ppm.

The carbonylation apparatus or process that is the subject of the invention typically includes a reactive section, purification section, a catalyst reservoir system and a light ends recovery system. The present invention may be 20 appreciated in connection with, for example, the carbonylation of methanol with carbon monoxide in a homogeneous catalytic reaction system comprising a reaction solvent (typically acetic acid), methanol and/or its reactive derivatives, a soluble rhodium catalyst, at least a finite concentration of water. The carbonylation reaction proceeds as methanol and carbon monoxide are 25 continuously fed to the reactor. The carbon monoxide reactant may be essentially pure or may contain inert impurities such as carbon dioxide, methane, nitrogen, noble gases, water and C₁ to C₄ paraffinic hydrocarbons. The presence of hydrogen in the carbon monoxide and generated *in situ* by the water gas shift reaction is preferably kept low, for example, less than 1 Bar partial pressure, as its 30 presence may result in the formation of hydrogenation products. The partial

pressure of carbon monoxide in the reaction is suitably in the range 1 to 70 bar, preferably 1 to 35 bar, and most preferably 1 to 15 bar.

5 The pressure of the carbonylation reaction is suitably in the range 10 to 200 Bar, preferably 10 to 100 bar, most preferably 15 to 50 Bar. The temperature of the carbonylation reaction is suitably in the range 100 to 300°C, preferably in the range 150 to 220°C. Acetic acid is typically manufactured in a liquid phase reaction at a temperature of from about 150 - 200°C and a total pressure of from about 20 to about 50 bar.

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Acetic acid is typically included in the reaction mixture as the solvent for the reaction.

15 Suitable reactive derivatives of methanol include methyl acetate, dimethyl ether, methyl formate and methyl iodide. A mixture of methanol and reactive derivatives thereof may be used as reactants in the process of the present invention. Preferably, methanol and/or methyl acetate are used as reactants. At least some of the methanol and/or reactive derivative thereof will be converted to, and hence present as, methyl acetate in the liquid reaction composition by reaction
20 with acetic acid product or solvent. The concentration in the liquid reaction composition of methyl acetate is suitably in the range 0.5 to 70% by weight, preferably 0.5 to 50% by weight, more preferably 1 to 35% by weight and most preferably 1-20% by weight.

25 Water may be formed *in situ* in the liquid reaction composition, for example, by the esterification reaction between methanol reactant and acetic acid product. Water may be introduced to the carbonylation reactor together with or separately from other components of the liquid reaction composition. Water may be separated from other components of reaction composition withdrawn from the
30 reactor and may be recycled in controlled amounts to maintain the required concentration of water in the liquid reaction composition. Preferably, the

concentration of water maintained in the liquid reaction composition is in the range 0.1 to 16% by weight, more preferably 1 to 14% by weight, most preferably 1 to 10% by weight.

5 The reaction liquid is typically drawn from the reactor and flashed in a one step or multi-step process using a converter as well as a flash vessel as hereinafter described. The crude vapor process stream from the flasher is sent to a purification system which generally includes at least a light ends column and a dehydration column. As noted earlier, the form of any catalyst metal which is
10 entrained to the light ends column and beyond is not well understood; however, the entrained catalyst metal is lost in conventional systems.

 The present invention is further appreciated by reference to **Fig. 1** which is a schematic diagram illustrating a typical carbonylation process and apparatus. In
15 **Fig. 1** there is shown a carbonylation system **10** including a reactor **12** provided with a feed system **14** including a methanol surge tank **16** and a carbon monoxide feed line. A catalyst reservoir system includes a methyl iodide storage vessel **20** as well as a catalyst storage tank **22**. Reactor **12** is provided with a vent **24** and an optional vent **24a**. Reactor **12** is coupled to a flash vessel **26** by way of a conduit
20 **28** and optionally by way of vent **24a**. The flasher, in turn, is coupled to a purification section **30** which includes a light ends or stripper column **32**, a dehydration column **34** and a strong acid, silver-exchanged cation ion-exchange resin bed **36** which removes iodides from the product. Instead of a silver-exchanged, strong acid cation ion-exchange resin, it has been reported that anion
25 ion-exchange resin can be used to remove iodides. *See* British Patent No. G 2112394A, as well as United States Patent No. 5,416,237, Col. 7, lines 54+, which teaches the use of 4-vinylpyridine resins for iodide removal.

 A gaseous purge stream is typically vented from the head of the reactor to
30 prevent buildup of gaseous by-products such as methane, carbon dioxide and hydrogen and to maintain a set carbon monoxide partial pressure at a given total

reactor pressure. A very significant improvement in processing involves minimizing or eliminating the high pressure vent from reactor 12 via line 24 to a high pressure absorber and instead utilizing a vent line such as line 24a. When operating at low water conditions as described herein, by-products and vent requirements are much reduced such that non-condensables can be vented at low pressure after flashing and stripping the light ends as is seen in Figures 1 and 2 while maintaining a predetermined carbon monoxide partial pressure in the reactor. Thus, the use of a high pressure absorber can be eliminated and/or minimized saving capital and operating costs.

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Optionally (as illustrated in Chinese Patent No. ZL92108244.4), a so-called "converter" reactor can be employed which is located between the reactor and flasher vessel shown in Figure 1. Optionally, the gaseous purge streams may be vented through the flasher base liquid or lower part of the light ends column to enhance rhodium stability and/or they may be combined with other gaseous process vents (such as the purification column overhead receiver vents) prior to scrubbing. Carbon monoxide may be added directly to a converter vessel if so desired or may be added slightly before (upstream) or after (downstream) if so desired in order to stabilize the catalyst solution and consume any unreacted methanol. Details of such arrangements are seen in European Patent No. EP 0 759 419 as well as United States Patent No. 5,770, 768 to *Denis et al.*, the disclosures of which are hereby incorporated by reference.

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These variations are well within the scope of the present invention as will be appreciated from the appended claims and the description which follows.

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As will be appreciated by one of skill in the art, the different chemical environments encountered in the purification train may require different metallurgy. For example, equipment at the outlet of the light ends column will likely require a zirconium vessel due to the corrosive nature of the process stream,

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while a vessel of stainless steel may be sufficient for equipment placed downstream of the dehydration column where conditions are much less corrosive.

Carbon monoxide and methanol are introduced continuously into reactor 12 with adequate mixing at a high carbon monoxide partial pressure. The non-condensable bi-products are vented from the reactor to maintain an optimum carbon monoxide partial pressure. The reactor off gas is treated to recover reactor condensables, *i.e.*, methyl iodide before flaring. Methanol and carbon monoxide efficiencies are generally greater than about 98 and 90% respectively. As will be appreciated from the *Smith et al.* patent noted above, major inefficiencies of the process are the concurrent manufacture of carbon dioxide and hydrogen by way of the water gas shift reaction.

From the reactor, a stream of the reaction mixture is continuously fed via conduit 28 to flasher 26. Through the flasher the product acetic acid and the majority of the light ends (methyl iodide, methyl acetate, water) are separated from the reactor catalyst solution, and the crude process stream 38 is forwarded with dissolved gases to the distillation or purification section 30 in single stage flash. The catalyst solution is recycled to the reactor via conduit 40.

The purification of the acetic acid typically includes distillation in a light ends column, a dehydration column, and, optionally, a heavy ends column. The crude vapor process stream 38 from the flasher is fed into the light ends column 32. Methyl iodide, methyl acetate, and a portion of the water condense overhead in the light end columns to form two phases (organic and aqueous) in a receiver 42. Both overhead liquid phases return to the reaction section via recycle line 44. Optionally, a liquid recycle stream 45 from the light ends column may also be returned to the reactor.

The purified process stream 50 is drawn off the side of the light ends column 32 and is fed into dehydration column 34. Water and some acetic acid

from this column separate and are recycled to the reaction system via recycle line 44 as shown. The purified and dried process stream 52 from the dehydration column 34 feeds resin bed 36 and product is taken therefrom at 56 as shown. Carbonylation system 10 uses only two primary purification columns and is
5 preferably operated as described in more detail in United States Patent No. 6,657,078 to *Scates et al.*, entitled "Low Energy Carbonylation Process", the disclosure of which is incorporated herein by reference. Additional columns are generally used as desired, depending on the system.

10 Receiver 42 is vented via line 60 to the light ends recovery system 70 shown in **Figure 2** which includes a switching system 72 which has a plurality of valves and pumps in order to selectively couple system 70 to scrubber solvent sources and return the used scrub solvent to the desired point in the carbonylation system as hereinafter described. Note also reactor 12 may be directly vented to
15 system 70 via line 24 if necessary.

Light ends recovery system 70 has an absorber tower 75 which is fed with vent gas via line 80 and with scrubber solvent via line 82. Preferably the scrubber solvent is chilled with a chiller 84 prior to being fed to tower 75 wherein the
20 solvent flows countercurrently with respect to the vent gas, absorbing methyl iodide and additional relative components before exiting the tower via return line 84 and being returned to the carbonylation unit. The scrubbed vent gas exits the tower at 86 and is further processed. For example, a second stage water scrub could be used to remove methyl acetate, methanol, acetic acid and so forth before
25 flaring if so desired. Alternatively, a second stage water scrub could be provided in tower 75 if so desired. Preferably, more than 90% of the methyl iodide is removed from the vent gas. The scrubber fluid is generally chilled to a temperature of from about 5°C to about 25°C prior to use in the tower, with the proviso that when acetic acid is used as the scrubber solvent, the temperature of
30 the solvent is held at 17°C or more to prevent freezing.

Switching system 72 includes a plurality of valves such as valves 90, 92, 94, 96, 98 and one or more pumps 100, 102 to raise pressure in the return lines 104, 106, 108, 110 if needed. Feed valves 96, 98 are used to select the scrubber solvent which may be methanol from tank 16 or acetic acid from stream 56
5 depending upon the mode of operation of tower 75.

In steady state operation of carbonylation system 10 valve 98 is closed and methanol is fed from tank 16 through open valve 96 via line 112 to chiller 84, wherein the methanol is cooled. From the chiller, methanol is fed to tower 75,
10 where it flows countercurrently with vent gas and sorbs methyl iodide and other volatile components therefrom before exiting the column via line 84. The used solvent with sorbed methyl iodide is pumped back to reactor 12 or tank 16 with pumps 100, 102 via line 106. In this mode of operation valves 92, 94 are closed and valve 90 is open.

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During start up or shut down of the system it may be desirable to operate tower 75 using acetic acid as the scrub solvent. In this mode of operation, valve 98 is open and valve 96 is closed. Acid may be sourced from product stream 56 or a tank from (TF) if so desired. The acid flows through line 112 to chiller 84
20 where it is chilled and fed to tower 75 via line 82 and scrubs the vent gas supplied via lines 60, 80 as noted above. The acid exits the tower 75 via line 84 and is pumped back to the carbonylation system by way of pumps 100, 102 via lines 104, 108. In this mode of operation of tower 75, valves 90, 94 are closed and valve 92 is open so that the used acetic acid is returned to light ends column 32,
25 the dehydration column 34, or elsewhere in the purification system for stripping.

During changeover from one solvent to the other, such as from methanol to acetic acid, it is generally undesirable to return the scrub fluid to the methanol feed system or light ends column since inefficiencies result. For such, a
30 changeover may be accomplished in from about 5 to about 20 minutes, during which time the used scrubber solvent is fed to catalyst reservoir 22. In changeover

mode, valves 90, 92 are closed and valve 94 is open. Thus the system is operated generally by way of (a) feeding vent gas from the carbonylation unit to the absorber tower, the vent gas including methyl iodide and optionally additional volatile components; (b) supplying a first scrubber solvent to the absorber tower, the first scrubber solvent consisting essentially of acetic acid; (c) contacting the vent gas with the first scrubber solvent thereby removing methyl iodide and optionally additional volatile components from the gas and absorbing methyl iodide and optionally additional volatile components into the first scrubber solvent; (d) feeding an absorber return stream including first scrubber solvent and absorbed methyl iodide and optionally additional absorbed volatile components to the light ends column, the dehydration column or elsewhere in the purification system; (e) terminating the supply of first scrubber solvent to the absorber tower; (f) supplying a second scrubber solvent to the absorber tower, the second scrubber solvent consisting essentially of methanol; (g) contacting the vent gas with the second scrubber solvent thereby removing methyl iodide and optionally additional volatile components from the gas and absorbing methyl iodide and optionally additional volatile components into the second scrubber solvent; (h) feeding an absorber return stream including first scrubber solvent, second scrubber solvent, absorbed methyl iodide and optionally additional absorbed volatile components from the absorber tower to the reactor; and (i) following the transition period, continue feeding an absorber return stream including second scrubber solvent and absorbed methyl iodide and optionally additional absorbed volatile components to the reactor. Feed to the absorber tower is selected by operation of valves 96, 98.

While the invention has been illustrated in connection with a particular apparatus, modifications to these examples within the spirit and scope of the invention will be readily apparent to those of skill in the art. In view of the foregoing discussion, relevant knowledge in the art and references discussed above in connection with the Background and Detailed Description, the disclosures of which are all incorporated herein by reference, further description is deemed unnecessary.

WHAT IS CLAIMED IS:

1. An apparatus for producing acetic acid comprising:
 - 5 (a) a reactor for carbonylating methanol or its reactive derivatives, the reactor containing a catalyst selected from rhodium catalysts, iridium catalysts and mixtures thereof, and a methyl iodide promoter in an acetic acid reaction mixture;
 - 10 (b) a feed system for providing carbon monoxide and methanol or its reactive derivatives to the reactor;
 - (c) a flash system configured to receive a stream of the reaction mixture and separate it into (i) at least a first liquid recycle stream, and (ii) a crude process stream containing acetic acid;
 - 15 (d) a first distillation column coupled to the flash system, the first distillation column being adapted to separate low boiling components including methyl iodide from the crude process stream, and generate a purified process stream,
 - 20 the first distillation column and optionally the reactor and flash system also operating to generate a vent gas stream comprising volatile organic components including methyl iodide;
 - 25 (e) an absorber tower adapted for receiving the vent gas stream and removing methyl iodide therefrom with a scrubber solvent, the absorber tower also being coupled to first and second scrubber solvent sources which are capable of supplying different first and second scrubber solvents; and
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(f) a switching system for alternatively providing first or second scrubber solvents to the absorber tower from either the first scrubber solvent source or the second scrubber solvent source.

- 5 2. The apparatus according to Claim 1, wherein the first scrubber solvent comprises methanol and the second scrubber solvent consists essentially of acetic acid.
3. The apparatus according to Claim 2, wherein the first scrubber solvent consists
10 essentially of methanol.
4. The apparatus according to Claim 1, further comprising a chiller coupled to the absorber tower and first and second scrubber solvent sources for cooling the scrubber solvents.
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5. The apparatus according to Claim 1, wherein a return stream from the absorber tower is selectively coupled to the feed system to the reactor or the first and/or second distillation columns.
- 20 6. The apparatus according to Claim 1, further including a catalyst reservoir system and wherein a return stream from the absorber tower is selectively coupled to the feed system to the reactor, the first and/or second distillation columns, or the catalyst reservoir system.
- 25 7. The apparatus according to Claim 1, wherein the feed system includes a methanol surge tank which is connected to the absorber tower as the first scrubber solvent source.
8. The apparatus according to Claim 1, further comprising a drying column
30 coupled to said first distillation column adapted for receiving the purified product stream and removing water therefrom.

9. The apparatus according to Claim 1, wherein the reactor is vented to the flash system.
- 5 10. A method of operating an absorber tower in a carbonylation unit for making acetic acid of the class including a reactor and a light ends column comprising:
- 10 (a) feeding vent gas from the carbonylation unit to the absorber tower, the vent gas including methyl iodide and optionally additional volatile components;
- (b) supplying a first scrubber solvent to the absorber tower, the first scrubber solvent consisting essentially of acetic acid;
- 15 (c) contacting the vent gas with the first scrubber solvent thereby removing methyl iodide and optionally additional volatile components from the gas and absorbing methyl iodide and optionally additional volatile components into the first scrubber solvent;
- 20 (d) feeding an absorber return stream including first scrubber solvent and absorbed methyl iodide and optionally additional absorbed volatile components to the light ends column and/or dehydration column;
- (e) terminating the supply of first scrubber solvent to the absorber tower;
- 25 (f) supplying a second scrubber solvent to the absorber tower, the second scrubber solvent comprising methanol, methyl acetate or mixtures thereof;
- 30 (g) contacting the vent gas with the second scrubber solvent thereby removing methyl iodide and optionally additional volatile components

from the gas and absorbing methyl iodide and optionally additional volatile components into the second scrubber solvent; and

- 5 (h) feeding an absorber return stream including second scrubber solvent and absorbed methyl iodide and optionally additional absorbed volatile components to the reactor.

10 11. The method according to Claim 10, wherein said second scrubber solvent consists essentially of methanol.

12. The method according to Claim 10, further comprising the step of chilling the first scrubber solvent prior to supplying it to the absorber tower.

15 13. The method according to Claim 10, further comprising the step of chilling the second scrubber solvent prior to supplying it to the absorber tower.

20 14. The method according to Claim 10, further comprising mixing the second scrubber solvent and absorbed methyl iodide and optional additional absorbed volatile components with methanol or a reactive derivative thereof prior to feeding the used solvent to the reactor.

25 15. The method according to Claim 10, wherein the vent gas includes methyl acetate and the first and second scrubber solvents are effective to absorb methyl acetate from the vent gas.

16. A method of operating an absorber tower in a carbonylation unit for making acetic acid of the class including a reactor and a light ends column comprising:

- 30 (a) feeding vent gas from the carbonylation unit to the absorber tower, the vent gas including methyl iodide and optionally additional volatile components;

- (b) supplying a first scrubber solvent to the absorber tower, the first scrubber solvent consisting essentially of acetic acid;
- 5 (c) contacting the vent gas with the first scrubber solvent thereby removing methyl iodide and optionally additional volatile components from the gas and absorbing methyl iodide and optionally additional volatile components into the first scrubber solvent;
- 10 (d) feeding an absorber return stream including first scrubber solvent and absorbed methyl iodide and optionally additional absorbed volatile components to the light ends column and/or dehydration column;
- (e) terminating the supply of first scrubber solvent to the absorber tower;
- 15 (f) supplying a second scrubber solvent to the absorber tower, the second scrubber solvent consisting essentially of methanol;
- (g) contacting the vent gas with the second scrubber solvent thereby
- 20 removing methyl iodide and optionally additional volatile components from the gas and absorbing methyl iodide and optionally additional volatile components into the second scrubber solvent;
- (h) feeding an absorber return stream including first scrubber solvent,
- 25 second scrubber solvent, absorbed methyl iodide and optionally additional absorbed volatile components from the absorber tower to the reactor during a transition period; and
- (i) following the transition period, continue feeding an absorber return
- 30 stream including second scrubber solvent and absorbed methyl iodide and optionally additional absorbed volatile components to the reactor.

17. The method according to Claim 16, wherein said second scrubber solvent consists of methanol.

5 18. The method according to Claim 16, further comprising the step of chilling the first scrubber solvent prior to supplying it to the absorber tower.

19. The method according to Claim 16, further comprising the step of chilling the second scrubber solvent prior to supplying it to the absorber tower.

10

20. The method according to Claim 16, further comprising mixing the second scrubber solvent and absorbed methyl iodide and optional additional absorbed volatile components with methanol or a reactive derivative thereof prior to feeding the used solvent to the reactor.

15

21. An improved method of operating an apparatus for making acetic acid of the class including a production system having a reactor containing a reaction medium and a product purification train wherein the production system is vented to an absorber tower, the improvement comprising:

20

(a) carbonylating methanol or reactive derivatives thereof in the reactor in the presence of a Group VIII metal catalyst and a methyl iodide promoter while maintaining a concentration of water in the reactor of from 1-10% by weight of the reaction medium and concurrently
25 maintaining in the reactor a predetermined partial pressure of carbon monoxide;

25

(b) venting non-condensables from the production system so as to provide low pressure vent gas only;

30

- (c) feeding the low pressure vent gas from the production system to the absorber tower, the vent gas including methyl iodide and optionally additional volatile components;
- 5 (d) supplying a scrubber solvent to the absorber tower, the scrubber solvent comprising methanol, methyl acetate or mixtures thereof;
- (e) contacting the low pressure vent gas with the scrubber solvent thereby removing methyl iodide and optionally additional volatile components
- 10 from the gas and absorbing methyl iodide and optionally additional volatile components into the scrubber solvent; and
- (f) feeding an absorber return stream including scrubber solvent and absorbed methyl iodide and optionally additional absorbed volatile
- 15 components to the reactor.
22. The method according to Claim 21, wherein the Group VIII metal catalyst is a supported catalyst.
- 20 23. The method according to Claim 22, wherein the Group VIII metal catalyst is a supported rhodium catalyst.
24. The method according to Claim 23, wherein the rhodium catalyst is supported on a crosslinked polyvinylpyrrolidone polymer.
- 25 25. The method according to Claim 23, wherein the rhodium catalyst is supported on a crosslinked polyvinylpyridine polymer.

FIG. 1

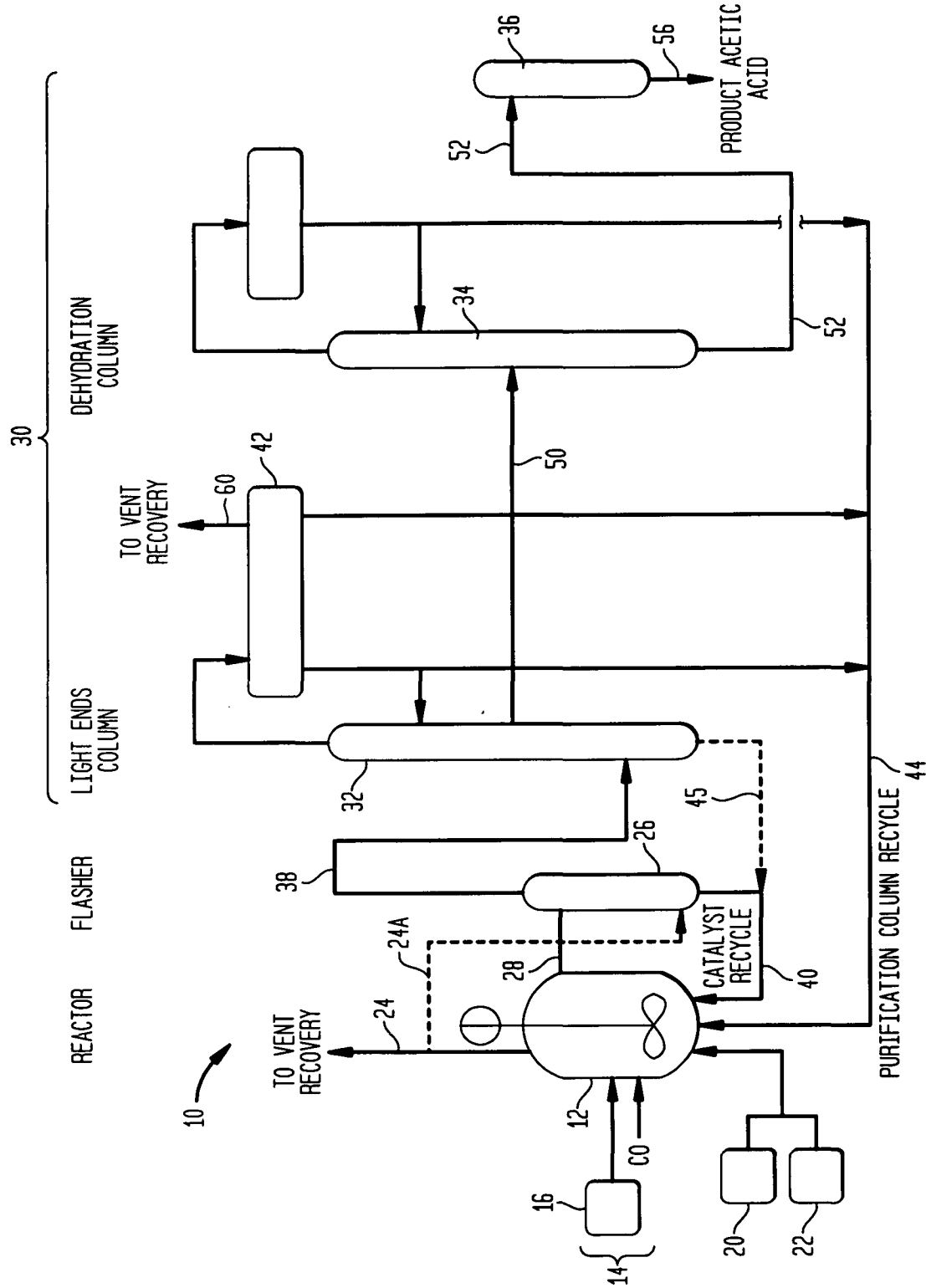


FIG. 2

