Title: INTEGRATED METHOD FOR THE PRODUCTION OF TRIOXANE FROM FORMALDEHYDE

Abstract:
The invention relates to an integrated method for producing trioxane from formaldehyde. According to said method, a flow A1 containing water and formaldehyde as well as a recirculation flow B2 substantially composed of water and formaldehyde are delivered to a trioxane synthesis reactor in which the formaldehyde is reacted to trioxane such that a product flow A2 containing trioxane, water, and formaldehyde is obtained. Flow A2 and a recirculation flow D1 containing trioxane, water, and formaldehyde are fed to a first distillation column and are distilled at a pressure ranging from 0.1 to 2.5 bar such that a trioxane-enriched flow B1 and flow B2, which is essentially composed of water and formaldehyde, are obtained. Flow B1 is delivered to a second distillation column and is distilled at a pressure ranging from 0.2 to 17.5 bar such that a product flow substantially comprising trioxane and a flow C1 containing trioxane, water, and formaldehyde are obtained. Flow C1 is fed to a third distillation column and is distilled at a pressure ranging from 1 to 10 bar such that the recirculation flow D1 containing trioxane, water, and formaldehyde as well as a flow D2 that is substantially composed of water are obtained.
Title: INTEGRATED METHOD FOR THE PRODUCTION OF TRIOXANE FROM FORMALDEHYDE

Bezeichnung: INTEGRIERTES VERFAHREN ZUR HERSTELLUNG VON TRIOXAN AUS FORMALDEHYD

Abstract: The invention relates to an integrated method for producing trioxane from formaldehyde. According to a said method, a flow A containing water and formaldehyde as well as a recirculation flow B2 substantially composed of water and formaldehyde are delivered to a trioxane synthesis reactor in which the formaldehyde is reacted to trioxane such that a product flow A2 containing trioxane, water, and formaldehyde is obtained. Flow A2 and a recirculation flow D1 containing trioxane, water, and formaldehyde are fed to a first distillation column and are distilled at a pressure ranging from 0.1 to 2.5 bar such that a trioxane-enriched flow B1 and flow B2, which is essentially composed of water and formaldehyde, are obtained. Flow B1 is delivered to a second distillation column and is distilled at a pressure ranging from 0.2 to 17.5 bar such that a product flow substantially comprising trioxane and a flow C1 containing trioxane, water, and formaldehyde are obtained. Flow C1 is fed to a third distillation column and is distilled at a pressure ranging from 1 to 10 bar such that the recirculation flow D1 containing trioxane, water, and formaldehyde as well as a flow D2 that is substantially composed of water are obtained.

Zusammenfassung: Die Erfindung betrifft ein integriertes Verfahren zur Herstellung von Trioxan aus Formaldehyd, bei dessen in einem ersten Schritt ein Wasser und Formaldehyd enthaltender Strom A1 und ein im Wesentlichen aus Wasser und Formaldehyd bestehender Rückführstrom B2 einem Trioxan-Synthesereaktor zugeführt werden, in welchem das Formaldehyd zu Trioxan umgesetzt wird, wobei ein Trioxan, Wasser und Formaldehyd enthaltender Produktstrom A2 erhalten wird. Der Strom A2 und ein Trioxan, Wasser und Formaldehyd enthaltender Rückführstrom D1 werden einer ersten Destillationskolonne zugeführt und bei einem Druck im Bereich von 0,1 bis 2,5 bar destilliert, wobei ein an Trioxan angereicherter Strom B1 und der im Wesentlichen aus Wasser und Formaldehyd bestehende Strom B2 erhalten werden. Der Strom B1 wird einer zweiten Destillationskolonne zugeführt und bei einem Druck im Bereich von 0,2 bis 17,5 bar destilliert, wobei ein im Wesentlichen aus Trioxan bestehender Produktstrom C2 und ein Trioxan, Wasser und Formaldehyd enthaltender Strom C1 erhalten werden. Der Strom C1 wird einer dritten Destillationskolonne zugeführt und bei einem Druck im Bereich von 1 bis 10 bar destilliert, wobei der Trioxan, Wasser und Formaldehyd enthaltende Rückführstrom D1 und ein im Wesentlichen aus Wasser bestehender Strom D2 erhalten werden.
Zur Erklärung der Zweitbuchstaben-Codes und der anderen Abkürzungen wird auf die Erklärungen ("Guidance Notes on Codes and Abbreviations") am Anfang jeder regulären Ausgabe der PCT-Gazette verwiesen.
INTEGRATED METHOD FOR THE PRODUCTION OF TRIOXANE FROM FORMALDEHYDE

Description

The invention relates to an integrated process for preparing trioxane from formaldehyde.

Trioxane is generally prepared by reactive distillation of aqueous formaldehyde solution in the presence of acidic catalysts. This affords a mixture comprising trioxane, formaldehyde and water as distillate. The trioxane is subsequently extracted from this mixture by extraction with halogenated hydrocarbons such as methylene chloride or 1,2-dichloroethane, or other water-immiscible solvents.

DE-A 1 668 867 describes a process for removing trioxane from mixtures comprising water, formaldehyde and trioxane by extraction with an organic solvent. In this process, an extraction zone consisting of two subzones is charged at one end with an organic, virtually water-immiscible extractant for trioxane, and at the other end with water. Between the two subzones, the distillate from the trioxane synthesis to be separated is fed. On the side of the solvent feed, an aqueous formaldehyde solution is then obtained, and, on the side of the water feed, a virtually formaldehyde-free solution of trioxane in the solvent.

A disadvantage of this procedure is the occurrence of extractant which has to be purified. Some of the extractants used are hazardous substances (T or T* substances in the context of the German Hazardous Substances Directive), whose handling entails special precautions.

DE-A 197 32 291 describes a process for removing trioxane from an aqueous mixture which consists substantially of trioxane, water and formaldehyde, by removing trioxane from the mixture by pervaporation and separating the trioxane-enriched permeate by rectification into pure trioxane on the one hand and an azeotropic mixture of trioxane, water and formaldehyde on the other. In one example, an aqueous mixture consisting of 40% by weight of trioxane, 40% by weight of water and 20% by weight of formaldehyde is separated in a first
distillation column under standard pressure into a water/formaldehyde mixture and into an azeotropic trioxane/water/formaldehyde mixture. The azeotropic mixture is passed into a pervaporation unit which comprises a membrane composed of polydimethylsiloxane with a hydrophobic zeolite. The trioxane-enriched mixture is separated in a second distillation column under standard pressure into trioxane and, in turn, into an azeotropic mixture of trioxane, water and formaldehyde. This azeotropic mixture is recycled upstream of the pervaporation stage.

This procedure is very costly and inconvenient. The pervaporation unit in particular entails high capital costs.

It is an object of the invention to provide an alternative process for preparing trioxane from aqueous formaldehyde solution to obtain pure trioxane. It is a particular object to provide a process which avoids the performance of extraction steps or pervaporation steps for obtaining pure trioxane.

The object is achieved by an integrated process for preparing trioxane from formaldehyde, which comprises the following steps:

a) a stream A1 comprising water and formaldehyde and a recycle stream B2 consisting substantially of water and formaldehyde are fed to a trioxane synthesis reactor in which the formaldehyde is converted to trioxane to obtain a product stream A2 comprising trioxane, water and formaldehyde;

b) stream A2 and a recycle stream D1 comprising trioxane, water and formaldehyde are fed to a first distillation column and distilled at a pressure in the range from 0.1 to 2.5 bar to obtain a stream B1 enriched in trioxane, and the stream B2 consisting substantially of water and formaldehyde;

c) stream B1 is fed to a second distillation column and distilled at a pressure in the range from 0.2 to 17.5 bar to obtain a product stream C2 consisting substantially of trioxane, and a stream C1 comprising trioxane, water and formaldehyde;

d) stream C1 is fed to a third distillation column and distilled at a pressure in the range from 1 to 10 bar to obtain the recycle stream D1 comprising
trioxane, water and formaldehyde, and a stream D2 consisting substantially of water.

Consisting substantially of one or more components means that these components are present to an extent of at least 90% by weight, preferably to an extent of at least 95% by weight, in the appropriate stream.

It is known that trioxane, formaldehyde and water form a ternary azeotrope which, at a pressure of 1 bar, has the composition of 69% by weight of trioxane, 5% by weight of formaldehyde and 26% by weight of water. According to the invention, the ternary azeotrope is separated by a pressure swing distillation, by carrying out a first and a second distillation stage at different pressures. In a first distillation stage which is operated at lower pressure, the starting mixture is separated into a trioxane-rich trioxane/water/formaldehyde mixture with low formaldehyde content on the one hand and a substantially trioxane-free formaldehyde/water mixture on the other. The trioxane-rich trioxane/water/formaldehyde mixture is subsequently separated in a second distillation stage which is carried out at high pressure into a trioxane-rich trioxane/water/formaldehyde mixture on the one hand and substantially pure trioxane on the other. According to the invention, the trioxane-rich trioxane/water/formaldehyde mixture is fed to a third distillation stage which is operated at a pressure which is between the pressure of the first distillation stage and of the second distillation stage. In the third distillation stage, a stream comprising trioxane, water and formaldehyde and a stream consisting substantially of water are obtained. The stream comprising trioxane, water and formaldehyde is recycled into the first distillation stage. This achieves substantially all trioxane prepared in the synthesis being obtained as a product of value.

According to the invention, each distillation stage comprises a distillation column. Suitable distillation columns are any distillation columns such as columns with structured packing or random packing. The distillation columns may comprise any internals, structured packings or random packings. In the following, all pressure data relate to the pressure at the top of the column in question.

In a first process step a), a stream A1 comprising water and formaldehyde and a recycle stream B2 consisting substantially of water and formaldehyde are fed to a trioxane synthesis reactor and allowed to react to obtain a product stream A2
comprising trioxane, water and formaldehyde. The reaction is effected under acidic conditions.

Streams A1 and B2 can be fed separately. However, it is also possible to mix streams A1 and B2 before they are fed into the trioxane synthesis reactor.

In general, stream A1 comprises from 50 to 85% by weight of formaldehyde and from 15 to 50% by weight of water.

The ratio of streams A1 and B2 is preferably selected such that, overall, from 15 to 70% by weight of water and from 30 to 85% by weight of formaldehyde, more preferably from 20 to 63% by weight of water and from 37 to 80% by weight of formaldehyde, are fed to the trioxane synthesis reactor.

Product stream A2 comprises generally from 35 to 84% by weight of formaldehyde, from 15 to 45% by weight of water and from 1 to 30% by weight of trioxane.

In one embodiment of the process according to the invention, the water/formaldehyde mixture is reacted in the trioxane synthesis stage a) in the presence of acidic homogeneous or heterogeneous catalysts such as ion exchange resins, zeolites, sulfuric acid and p-toluenesulfonic acid at a temperature of generally from 70 to 130°C. It is possible to work in a reactive distillation column or a reactive evaporator. The product mixture composed of trioxane, formaldehyde and water is then obtained as a vaporous vapor draw stream of the reactive evaporator or as a top draw stream of the reactive distillation column. The trioxane synthesis may also be carried out in a fixed bed reactor or fluidized bed reactor over a heterogeneous catalyst, for example an ion exchange resin or zeolite.

In a step b) which follows step a), stream A2 and a recycle stream D1 comprising trioxane, water and formaldehyde are fed to a first distillation column and distilled at a pressure of from 0.1 to 2.5 bar, preferably from 0.3 to 2 bar, more preferably from 0.4 to 1.5, for example 1 bar, to obtain a stream B1 enriched in trioxane, and the stream B2 consisting substantially of water and formaldehyde.
The first distillation column comprises preferably from 2 to 50, more preferably from 4 to 40 theoretical plates. In general, the stripping section of the first distillation column comprises at least 25% of the number of theoretical plates of the distillation column. The rectifying section preferably comprises from 50 to 90% of the theoretical plates of this distillation column.

The stream B1 enriched in trioxane comprises generally from 30 to 80% by weight of trioxane, from 20 to 69% by weight of water and from 1 to 20% by weight of formaldehyde. The trioxane-enriched stream B1 preferably comprises from 60 to 75% by weight of trioxane, from 5 to 35% by weight of water and from 5 to 20% by weight of formaldehyde. Stream B2 comprises generally from 51 to 85% by weight of formaldehyde, from 15 to 49% by weight of water and from 0 to 1% by weight of trioxane. Stream B2 preferably comprises less than 0.5% by weight of trioxane, more preferably less than 0.1% by weight of trioxane. Stream D1 comprises generally from 5 to 30% by weight of formaldehyde, from 5 to 30% by weight of water and from 50 to 80% by weight of trioxane. Stream D1 comprises preferably from 5 to 20% by weight of formaldehyde, from 10 to 40% by weight of water and from 55 to 70% by weight of trioxane.

Streams A2 and D1 are fed to the first distillation column preferably as side feeds. Stream B1 is withdrawn from the first distillation column preferably as a top draw stream and stream B2 as a bottom draw stream.

In a preferred embodiment stream A2 is fed to the first distillation column as a side feed in the stripping section or as a feed in the bottom of the column and the stream D1 as a side feed in the rectifying section. In a further embodiment the stream A2 and the stream D1 can be mixed before feeding in the first distillation column. The feeding then preferably is carried out as a combined side feed.

In a further embodiment of the process according to the invention, the trioxane synthesis stage a) and the first distillation stage b) are carried out together as a reactive distillation in a reaction column. In the stripping section, this may comprise a fixed catalyst bed of a heterogeneous catalyst. Alternatively, the reactive distillation may also be carried out in the presence of a homogeneous catalyst, in which case an acidic catalyst is present together with the water/formaldehyde mixture in the column bottom. Preference is given to carrying
out the reactive distillation in the presence of a homogeneous acidic catalyst which is present in the column bottom together with the aqueous formaldehyde solution.

In a process step c) which follows step b), the trioxane-enriched stream B1 is fed to a second distillation column and distilled at a pressure of from 0.2 to 17.5 bar to obtain a stream C2 consisting substantially of pure trioxane, and a stream C1 comprising trioxane, water and formaldehyde.

The second distillation column comprises generally at least 2 theoretical plates, preferably from 10 to 50 theoretical plates. In general, the stripping section of this distillation column comprises from 25 to 90%, preferably from 50 to 75%, of the theoretical plates of this column.

The pressure in the second distillation column is at least 0.1 bar higher than in the first distillation column. In general, this pressure difference is from 0.5 to 10 bar, preferably from 1 to 7 bar. The second distillation column is operated preferably at a pressure between 2 and 10 bar, more preferably at a pressure between 2 and 7 bar.

Product stream C2 comprises generally from 95 to 100% by weight, preferably from 99 to 100% by weight of trioxane, and from 0 to 5% by weight, preferably from 0 to 1% by weight of water. More preferably, the content of water in the product stream is < 0.1%. It may even be < 0.01%. Stream C1 comprises, for example, from 5 to 20% by weight of formaldehyde, from 15 to 35% by weight of water and from 50 to 75% by weight of trioxane.

Preferably, stream B1 is fed as a side feed, preferably as a side feed in the stripping section of the second distillation column, to the second distillation column. Stream C1 is preferably withdrawn from the second distillation column as a top draw stream and product stream C2 as a bottom draw stream.

In the step d) which follows step c), stream C1 is fed to a third distillation column and distilled at a pressure in the range from 1 to 10 bar to obtain the recycle stream D1 comprising trioxane, water and formaldehyde and a product stream D2 consisting substantially of water.
The third distillation stage is carried out preferably at a pressure between 1 and 5 bar.

The third distillation column has at least two theoretical plates, preferably from 10 to 50 theoretical plates. In general, the stripping section of this distillation column comprises from 25 to 90%, preferably from 50 to 75% of the theoretical plates of this column.

Stream C1 is preferably fed to the third distillation column as a side feed in the stripping section of the column. Recycle stream D1 is generally obtained at the top of the third distillation column and the stream D2 consisting substantially of water is obtained as a bottom draw stream or as a side draw stream in the stripping section of the column.

In a preferred embodiment, in a further step of the process for preparing trioxane from formaldehyde, a feed stream E1 comprising water and formaldehyde is fed to a formaldehyde concentration unit. Stream A1 as a formaldehyde-rich bottom stream and a low-formaldehyde stream E2 as a top or vapor draw stream are withdrawn from the concentration unit.

The concentration of the feed stream E1 comprising water and formaldehyde is generally carried out in a distillation column or in an evaporator. The concentration is preferably carried out in an evaporator, more preferably in a continuous evaporator. Suitable continuous evaporators are, for example, circulation evaporators, falling-film evaporators, helical tube evaporators or thin-film evaporators. Particular preference is given to using falling-film evaporators to concentrate the water/formaldehyde mixture. The falling-film evaporator is operated generally at a pressure of from 50 to 200 mbar and a temperature of from 40 to 75°C.

The concentration step can be carried out as described, for example, in DE-A 199 25 870.

The formaldehyde-enriched stream A1 obtained in the concentration is generally withdrawn as a bottom draw stream; the low-formaldehyde stream E2 is
withdrawn as a top or vapor draw stream.

When a distillation column is used for concentration, the feed stream E1 comprising water and formaldehyde is preferably fed as a side feed.

In a preferred embodiment, the low-formaldehyde stream E2 is fed to the third distillation column. Stream E2 is preferably fed to the third distillation column as a side feed at the top of the column. In a further embodiment, stream E2 and stream C1 can first be mixed and subsequently added to the third distillation column as a combined side feed.

In addition to water, formaldehyde and trioxane, streams A2, B1, C1 and D2 in particular may also comprise up to 15% by weight, generally from 1 to 10% by weight of low boilers. Typical low boilers which may be formed in the trioxane synthesis and the subsequent distillative separation are methyl formate, methylal, dimethoxydimethyl ether, trimethoxydimethyl ether, methanol, formic acid and also further hemiacetals and full acetalts. To remove these low boilers, it is optionally possible after the first distillation stage b) to carry out a further distillation stage (low boiler removal stage). In this case, the low boilers are removed preferably via the top of the low boiler removal column which is preferably operated at a pressure of from 1 to 3 bar. In general, the low boiler removal column has at least 5 theoretical plates, preferably from 15 to 50 theoretical plates. The stripping section of this column comprises preferably from 25 to 90% of the theoretical plates of this column. Stream B1 is fed to this low boiler removal column as a side feed, and the stream B1’ freed of the low boilers is generally obtained as a bottom draw stream. When the low boiler removal is carried out, stream B1’ is fed as stream B1 to the downstream second distillation column.

The pure trioxane obtained, whose purity may be > 99% by weight, preferably > 99.5% by weight or even > 99.8% by weight, is preferably used to prepare polyoxymethylene (POM), polyoxymethylene derivatives such as polyoxymethylene dimethyl ether (POMDME), and diaminodiphenylmethane (MDA).

The invention is described in detail below with reference to the drawing.

The sole figure shows a process flow diagram of the process according to the
invention.

An aqueous formaldehyde solution 1 (stream E1) is added to a concentration unit 2. The concentration unit 2 may be any distillation column, for example a tray column, column with random packing or column with structured packing or a continuous evaporator, for example a circulation evaporator, falling-film evaporator, helical tube evaporator or thin-film evaporator. The concentration unit 2 is preferably a falling-film evaporator. From the concentration unit 2, a formaldehyde-rich bottom draw stream 3 (stream A1) and a low-formaldehyde aqueous vapor stream as top draw stream 4 (stream E2) are obtained. The formaldehyde-rich bottom draw stream 3 is fed to a trioxane synthesis reactor 5. In the trioxane synthesis reactor 5, the aqueous formaldehyde solution is reacted in the presence of an acidic catalyst present in homogeneous or heterogeneous form to give trioxane.

From the trioxane synthesis reactor 5, a stream 6 comprising trioxane, formaldehyde and water (stream A2) is fed as a side feed to a first distillation column 7. In the first distillation column 7, stream 6 is separated into a trioxane-enriched stream 8 (stream B1) which is withdrawn from the first distillation column 7 as a top draw stream and a stream 9 which is obtained as a top draw and consists substantially of water and formaldehyde (stream B2). The stream 9 obtained at the bottom (stream B2) is recycled into the trioxane synthesis reactor 5.

The stream 8 obtained at the top of the first distillation column 7 (stream B1) is fed to a second distillation column 10. In the second distillation column 10, this is separated into a product stream 11 comprising substantially trioxane (stream C2) and a stream 12 comprising trioxane, water and formaldehyde (stream C1) which is drawn off at the top of the second distillation column 10. Stream 14 is fed via a side feed to a third distillation column 13. In the third distillation column 13, stream 12 is separated into a recycle stream 14 which comprises trioxane, formaldehyde and water and is obtained at the top (stream D1) and a stream 15 consisting substantially of water (stream D2) which is drawn off at the bottom of the third distillation column.

The recycle stream 14 (stream D1) is fed to the first distillation column 7. The
recycle stream 14 is fed via a side feed into the first distillation column 7.

In addition to the stream 12 (stream C1), the low-formaldehyde stream 4 (stream E2) drawn off at the top of the concentration unit 2 is fed to the third distillation column 13. It is preferably fed as a side feed at the top of the third distillation column 13.

Example

An aqueous formaldehyde solution 1 comprising 37% by weight of formaldehyde and 63% by weight of water is added to a concentration unit 2 designed as a falling-film evaporator. The falling-film evaporator is operated at a pressure of 100 mbar and a temperature of 50°C. A bottom draw stream 3 comprising 50% by weight of formaldehyde and 50% by weight of water is withdrawn from the bottom of the falling-film evaporator. The top draw stream 4 comprises 20% by weight of formaldehyde, the remainder is water.

The bottom draw stream 3 is fed to the trioxane synthesis reactor 5. The trioxane synthesis reactor is designed as a stirred tank reactor and is operated at a temperature of 108°C. The discharge stream 6 comprises 9% by weight of trioxane and 66% by weight of formaldehyde, the remainder is water.

The stream 6 is fed to the first distillation column 7 on the fifth tray. The stream 14 is recycled from the process into this distillation column 7 on the 20th tray. The first distillation column 7 is operated at a pressure of 1 bar. The temperature at the top is about 101°C, the temperature at the bottom is about 104°C. The column contains 30 trays. A stream 9 with a concentration of 80% by weight of formaldehyde and 20% by weight of water is withdrawn from the bottom of the first distillation column. A stream 8 with a concentration of 66% by weight of trioxane, 7% by weight of formaldehyde, and 27% by weight of water is withdrawn from the top of the column 7.

The stream 8 is fed to the second distillation column 10. The second distillation column 10 is operated at a pressure of 4 bar. The temperature at the top is about 142°C, the temperature at the bottom is about 166°C. The column has 40 trays,
stream 8 is fed to the 20th tray. A stream 12 comprising 64% by weight of trioxane, 8% by weight of formaldehyde and 28% by weight of water is withdrawn from the top of the column 10. The product stream 11 comprising more than 99% by weight of trioxane is withdrawn from the bottom.

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The top stream 12 is fed to the third distillation column 13 on the 24th tray. Likewise, the stream 4 from the falling-film evaporator is fed to the third distillation column on the 36th tray. The third distillation column 13 contains 48 trays and is operated at a pressure of 2.5 bar. The temperature at the top is about 127°C, the temperature at the bottom is about 131°C. The stream 15 is withdrawn from the bottom of this column. This stream comprises more than 98% by weight of water. The top stream 14 comprises 66% by weight of trioxane, 8% by weight of formaldehyde and 26% by weight of water. This stream 14 is recycled to the first distillation column 7.

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as enclosed to IPRP

1. An integrated process for preparing trioxane from formaldehyde, which comprises the following steps:

   a) A stream A1 comprising water and formaldehyde and a recycle stream B2 consisting substantially of water and formaldehyde are fed to a trioxane synthesis reactor in which the formaldehyde is converted to trioxane to obtain a product stream A2 comprising trioxane, water and formaldehyde;

   b) Stream A2 is fed as a side feed in the stripping section of a first distillation column and a recycle stream D1 comprising trioxane, water and formaldehyde is fed as a side feed in the rectifying section of the first distillation column and distilled at a pressure in the range from 0.1 to 2.5 bar to obtain a stream B1 enriched in trioxane, and the stream B2 consisting substantially of water and formaldehyde;

   c) Stream B1 is fed to a second distillation column and distilled at a pressure in the range from 0.2 to 17.5 bar to obtain a product stream C2 consisting substantially of trioxane, and a stream C1 comprising trioxane, water and formaldehyde;

   d) Stream C1 is fed to a third distillation column and distilled at a pressure in the range from 1 to 10 bar to obtain the recycle stream D1 comprising trioxane, water and formaldehyde, and a stream D2 consisting substantially of water.

2. The process according to claim 1, wherein the distillation b) is carried out at a pressure in the range from 0.4 to 1.5 bar, the distillation c) at a pressure in the range from 2 to 7 bar and the distillation d) at a pressure in the range from 1 to 5 bar.

3. The process according to claim 1 or 2, wherein stream B1 is withdrawn as a top draw stream and stream B2 as a bottom draw stream from the first
distillation column and stream A2 is fed as a feed in the bottom of the column or as a side feed and stream D1 as a side feed.

4. The process according to any of claims 1 to 3, wherein stream B1 is fed as a side feed to the second distillation column and stream C1 is withdrawn as a top draw stream and stream C2 as a bottom draw stream.

5. The process according to any of claims 1 to 4, wherein stream C1 is fed as a side feed to the third distillation column, and stream D1 is withdrawn as a top draw stream and stream D2 as a bottom draw stream.

6. The process according to any of claims 1 to 5, wherein, in a further step, a feed stream E1 comprising water and formaldehyde is fed to a formaldehyde concentration unit, stream A1 is withdrawn from the concentration unit as a formaldehyde-rich bottom stream and a low-formaldehyde stream E2 is withdrawn as a top or vapor draw stream.

7. The process according to claim 6, wherein stream E2 is fed to the third distillation column.

8. The process according to claim 6, wherein stream E2 is fed to the third distillation column as a side feed at the top of the column.

9. The process according to claim 6, wherein stream E2 and stream C1 are mixed before addition to the third distillation column and then fed as a combined side feed to the third distillation column.

10. The process according to claim 6, wherein the formaldehyde concentration unit is a distillation column.

11. The process according to claim 6, wherein the formaldehyde concentration unit is an evaporator.

12. The process according to claim 11, wherein the evaporator is a falling-film evaporator.