Title: BAKE TYPE AROMATISING COMPOSITIONS

Abstract: Process for the preparation of aromatising compositions suitable for the biogeneration of mixtures comprising compounds being able to develop or enhance typical baked type aroma upon heating of bakery products such as bread an the like. Such process comprises the bioconversion of at least two amino compounds selected from the group consisting of amino acids and peptides and at least one reducing sugar in the presence of a micro-organism selected in the group consisting of yeasts. Method for making a bakery product having an improved aroma.
BAKED TYPE AROMATISING COMPOSITIONS

The present invention relates to a process for the preparation of aromatising compositions, more particularly the biogeneration of mixtures comprising compounds being able to develop or enhance typical baked type aroma upon heating.

Fresh bread-like aroma is one of the most important criteria of the quality of bakery products and such flavour reflects the freshness of the product. For packaged, chilled and frozen bakery products such as pizza, bread-rolls, croissants, crusts for example, this flavour is often weak mainly because of processing/storage of these products and thus the overall quality of the final product after heating or baking is generally perceived as not close to that of traditional freshly made products.

The bread-like aroma is composed of a complex mixture of odorant compounds identified in the crust of baguette (Zehentbauer G. and Grosh W., 1998. Journal of Cereal Science 28, 81-92.) such as 2-acetyl-1-pyrroline, 2-acetyl-2-thiazoline and pyrazines (responsible for roasted note), furanones (responsible for caramel-like note), ketones and diketones (responsible for butty note) and aldehydes (responsible for malty note), for example.

Several methods are known to prepare bread-type aromatising compositions mainly based on Maillard reaction and/or fermentation.

US 4663168 discloses a method for preparing a heat-stable yeast fermented malt reaction flavour concentrate from a mixture of malt flour, yeast and fermentable sugar. The method comprises the steps of fermenting a mixture of malt flour by yeast, heating the fermented malt to produce the
flavour and inactivating the yeast. The flavour obtained is described as bread crust-like, nutty and toasted grain. JP 7242661 discloses the preparation of dimethylpyrrolidiny1-furanone and its use for imparting or increasing freshly baked or boiled flavour of bakery foods. WO 9933358 describes a process for preparing an aromatising composition containing 2-acetyl-2-thiazoline and its precursors involving bioconversion of a sulfur containing compound and an organic acid or its derivative in the presence of yeast, separation and recovery of the surpernatant. This composition is presented as being able to enhance the roasted notes of bakery products. EP 535713 discloses the use of carbomethoxy-2-pyrroline-1 as aromatising agent to impart cereal or bread-crust aroma to product in which it is incorporated. EP 937402 describes a composition for intensifying the browning and the aroma of baked goods and comprises ascorbic acid, an amino acid, a carbohydrate and a phosphate and optionally lactic acid. This composition is intended to be sprayed onto pre-baked goods and thus both improves the appearance (browning) and intensifies the aroma.

All the previous works already done mainly focused on roasted aroma and moreover the aromatising compositions obtained were not fully balanced in the different typical aromas and flavours responsible for the global baked aroma. Moreover some of the previous methods were based on pure chemical reaction that is not always desired for food products.

The main purpose of the present invention is to provide a new and natural route for the manufacture of a well balanced baked aroma precursors composition which generates this baked aroma upon heating.
To this end, the process for the preparation of an aromatising composition according to the present invention, comprises the bioconversion of at least two amino compounds selected from the group consisting of amino acids and peptides and at least one reducing sugar in the presence of a micro-organism selected in the group consisting of yeasts.

The reaction mixture can advantageously be then submitted to a separation step, for example by centrifugation, so as to remove the cells from the reaction mixture. The obtained supernatant is usable as an aromatising composition for baked aroma generation upon heating either directly in liquid form or after drying in powder form obtained by mild dehydration methods, such as freeze-drying for example.

The amino acids to be subjected to the bioconversion may be advantageously selected from the group consisting of arginine, citrulline, glutamine, ornithine and proline. In the case of the use of peptides, the preferred forms consist of di-peptides and/or tripeptides of such chosen amino acids.

The reducing sugar of the mixture may be a mono or oligosaccharide starting from mono to tetra-saccharide, preferably a mono-saccharide and more preferably a C5 or C6 mono-saccharide. Thus the most preferred reducing sugar may be selected from the group consisting of fructose, glucose and rhamnose.

The combination of such compounds incubated with a micro-organism selected in the group consisting of yeasts allows to obtain a mixture containing key aroma precursors which can generate a fully rich and well-balanced baked aroma upon heating.

Preferred micro-organism used for the bioconversion may be baker's yeast belonging to the genus Saccharomyces
cerevisiae for example in the form of a powder, an extract, a compressed form or a cream solution. However other kind of micro-organisms may be used, such as Saccharomyces bayanus, Candida versatilis or Debaromyces hansenii, for example. Preferably, the yeast is freshly prepared, or up to 8 days old, advantageously up to 4 days old and kept in the refrigerator.

Regarding the respective quantities of the two starting substrate products, they can be such that the molar ratio between the amino compounds and the reducing sugar is about 1:1 or up to about 1:10. The concentration of these substrates in the reaction medium may range from about 10 to 1000 mMol, preferably from about 20 to 300 mMol.

Generally a yeast solution for bioconversion is used with a dry matter of from about 10 to 30%, preferably about 20% and more preferably about 17%. The substrate quantity may be used as from 10 to 100 mmol for about 100 ml of yeast solution showing about 20% dry matter. However, such range may be adjusted according to the yeast and the substrates concerned.

The incubation of substrates with yeasts, in other word, the bioconversion may be carried out either under aerobic or anaerobic conditions during 2 to 48 hours, preferably 6 to 12 hours and at a pH of about 5 to 8, preferably about 7. The pH value of the medium may be controlled and may be kept constant throughout the bioconversion and may thus be performed by means of a pH-stat, for example. The temperature of the bioconversion may range from about 20 to 50°C, preferably around 30 to 35°C, and may be carried out under low, medium or high agitation conditions.

The reaction medium may be water or any buffer solution such as phosphate buffer systems, for example.
The reaction is initiated by the addition of the substrates to the medium containing the micro-organism; this addition of substrates may be carried out in one time or in several times. For example, the total amount of reducing sugar may be added in two parts, one half at time 0 and one half afterward during bioconversion, for example.

As already mentioned, the reaction mixture advantageously may be submitted after bioconversion to a separation step, so as to recover the liquid phase from the cells. This separation step may be carried out by centrifugation, decantation, filtration or ultrafiltration, for example. The liquid phase may either be used as such as an aromatising composition or dried into a powder using mild conditions by freeze drying, for example. The powdering step may be carried out with or without support material such as maltodextrine, cyclodextrine or starch, for example.

The aromatising composition obtained by the process according to the present invention is able to generate a rich baked aroma upon heating; either heated in its liquid or powder form. The typical flavour obtained upon heating of the present aromatising composition can be described after sensory analysis as a fully rich and well balanced freshly baked aroma and can thus be characterized as roasted, bread crust-like, caramel-like, buttery and somewhat honey or yeasty as well.

As previously exposed, the baked aroma is composed of a complexe mixture of odorants compounds among which we can cite aldehydes, ketones and diketones, furane derivatives and alkylpyrazines. The typical flavour obtained upon heating of the aromatising composition obtained according to the present process is mainly due to the presence of such kind of compounds. Hence, several furane derivatives
such as furaneol has been identified after heating the composition according to the invention. Five aldehydes were also identified and among them methylpropanal, 2-methylbutanal and 3-methylbutanal were the target molecules exhibiting malty notes. Among the ketones identified, we can cite diacetyl, 2,3-pentanedione responsible for buttery note of bread crust aroma; moreover 3-hydroxy-2-butanone and 1-hydroxy-2-propanone were also identified exhibiting buttery and caramel-like notes. Concerning the alkylpyrazines generated, the smell of all these molecules was decribed as nutty and roasted with nuances from one to each other. The detailed results concerning the nature and relative quantities of these compounds can be seen in figures 1 to 4.

The heat treatment that reveals the baked aroma of the aromatising composition may be carried out directly on the composition itself, either in liquid or in powder form. However the heat treatment may also be done through heating of a foodstuff containing the aromatising composition obtained by the process of the present invention. The conditions for the heat treatment may be the followings: 90 to 200°C during 5 to 360 minutes depending on the form - liquid or powder - of the aromatising composition and on its concentration in flavour precursors. As already said, the heating treatment upon which the bread aroma develops may also be carried out through the heating of the foodstuff in which the aromatising composition has been introduced. Hence, the aromatising composition obtained according to the present process may be introduced into a dough or into a dough based product and thus will generate and/or enhance the baked aroma upon the baking/heating of the product. The products that may be concerned by the use of the aromatising composition obtained according to the present process may be bread doughs, pizza doughs, cereal products, biscuit snacks and batter doughs, crackers doughs, wafer
doughs, croissants and the like, for example. However, the use of the present aromatising composition is not limited to dough based products and may be used for any kind of food products intended to be heated, reheated, baked, toasted or whose manufacture process comprises a heating step and for which a typical fresh baked aroma is desired. Among such product we can cite breakfast cereals, cereals bars, cereal based confectionnaries, beverages and pet foods, for example. All the products concerned by the use of the present aromatising composition may be fresh, shelf-stable, chilled or frozen products. Hence, the original aromatising composition before any heating treatment may also be added to the various constituents and ingredients to be heated or applied as a coating for example by pulverizing the liquid composition onto prebaked or unbaked bakery products, for example. Despite that, the aromatising composition obtained by the bioconversion process according to the present invention may also be heated as such in order to obtain a baked aroma composition that may be added in, sprayed or sprinkled onto food products.

Thus, the present invention also concerns a method for making a bakery product having an improved aroma comprising the steps of

- mixing flour, water, yeast and an aromatising composition obtained by bioconversion of at least two amino compounds selected from the group consisting of amino acids and peptides and at least one reducing sugar in the presence of a micro-organism selected in the group consisting of yeasts,
- kneading all ingredient in order to obtain a dough,
- ferment the dough and
- bake the dough.
Such a method allows to obtain full baked aroma or enhance the aroma of bakery products even in the case of short fermentation time.

In another embodiment, the present invention also concerns the use of the aromatising composition obtained by the present bioconversion process in non fermented/non yeasted doughs in order to impart to such products a typical baked aroma.

Indeed, for non yeast-leavened dough, the aroma profile lacks the typical baked aroma. Thus the invention refers to a method for making a bakery product having an improved aroma comprising the steps of

- mixing at least flour, water and a aromatising composition obtained by bioconversion of at least two amino compounds selected from the group consisting of amino acids and peptides and at least one reducing sugar in the presence of a micro-organism selected in the group consisting of yeasts,

- kneading all ingredient in order to obtain a dough

- baking the dough.

The product thus obtained, despite the lack of yeast and fermenting step, exhibits an enhanced aroma and flavour profile typical to fresh bakery products.

The present invention will now be illustrated by reference to the following examples.

**EXAMPLE 1 :**

Preparation of aromatising composition A1 (ornithine / glutamine / rhamnose)

Active dry baker’s yeast (20 g) was added to distilled water (100 ml) and left under stirring for 20 min until
complete hydration. The yeast suspension was then centrifuged for 15 min at 3000 x g and at 4°C. Supernatant (70-75 ml) was discarded and replaced by the same volume of fresh distilled water. After suspension, the yeast was ready to be used.

Yeast suspension (100 g) was introduced into a glass reactor fitted with 3 necks. The temperature was controlled at 30°C (oil bath) and the pH was adjusted to 7.5 with 2 M NaOH and kept constant all along the fermentation, using a pH-stat system. Ornithine (2 mmol), glutamine (2 mmol) and rhamnose (10 mmol) were added at time 0 and another portion of rhamnose (10 mmol) was added after 4 hours. After 6 hours incubation time, the reaction mixture was centrifuged for 15 min at 3000 x g and at 4°C to remove yeast cells. The supernatant was recovered as composition A1. A part of this composition was freeze-dried to composition A1F.

EXAMPLE 2:
Preparation of aromatising compositions A2 and A2F (arginine / citruline / fructose).
The procedure was the same as in example 1 excepted that the quantity of respective components was: 6 mmol of arginine, 6 mmol of citruline and twice 30 mmol of fructose.

EXAMPLE 3:
Preparation of aromatising compositions A3 and A3F (arginine / citruline / rhamnose).
The procedure was the same as in example 1 excepted that the quantity of respective components was: 2 mmol of arginine, 2 mmol of citrulline and twice 10 mmol of rhamnose.

EXAMPLE 4:
Preparation of aromatising compositions A4 and A4F (arginine / citrulline / glucose).
The procedure was the same as in example 1 excepted that the quantity of respective components was: 2 mmol of arginine, 2 mmol of citrulline and twice 10 mmol of glucose.

EXAMPLE 5:
Preparation of aromatising compositions A5 and A5F (ornithine / glutamine / glucose).
The procedure was the same as in example 1 excepted that the quantity of respective components was: 2 mmol of ornithine, 2 mmol of glutamine and twice 10 mmol of glucose.

EXAMPLE 6:
Preparation of aromatising compositions A6 and A6F (ornithine / glutamine / fructose).
The procedure was the same as in example 1 excepted that the quantity of respective components was: 2 mmol of ornithine, 2 mmol of glutamine and twice 10 mmol of fructose.
EXAMPLE 7:
Generation of baked aroma by heat treatment.

One ml of sample solutions A1 to A6 or 60 mg of freeze-dried powder A1F to A6F, prepared as described in examples 1 to 6 were introduced into a 4 ml vial. The latter was closed, heated for 30, 45 minutes or 2h30 at 100°C in a multi-block heater and rapidly cooled down in a bath of ice. The sample was ready for sniffing or for further GC analysis.

Concerning the sniffing test, the panelists were asked to describe the aroma quality and intensity.

The results of these sniffing tests are shown in the following Table 1.

Table I: Aroma generated after heat treatment of aqueous solutions at 100°C for 30 min.

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<th>Samples</th>
<th>Aroma description</th>
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<tr>
<td>A2</td>
<td>Roasted, honey, yeasty</td>
</tr>
<tr>
<td>A6</td>
<td>Honey, roasted</td>
</tr>
<tr>
<td>A3</td>
<td>Roasted (strong), bread crust, caramel-like</td>
</tr>
<tr>
<td>A1</td>
<td>Roasted (very strong), bread crust, caramel-like</td>
</tr>
<tr>
<td>A4</td>
<td>Roasted (weak), yeasty, off-notes</td>
</tr>
<tr>
<td>A5</td>
<td>Roasted (weak), yeasty, off-notes</td>
</tr>
</tbody>
</table>
Head space Gas Chromatography analysis

Equipment:

- Automated Headspace sampler, Agilent Technologies model HP7694 coupled to a gas chromatograph, model 6890, equipped with a mass spectrometry detector, model 5973N.
- Capillary column: INNOWax, length: 60 m, inner diameter: 0.25 mm, film thickness: 0.50 μm, Agilent Technologies 19091N-236.
- Oven, Heraeus T5042E

Method:

Analysis of powder:

Sixty mg of powder (AlF), obtained as described in Example 1 were introduced into a vial for automated headspace sampler. The vial was sealed and placed into the autosampler. Headspace analysis was performed without any shaking. Sample vial was heated in the oven set at 100°C for 5, 10, 15, 20, 30 or 45 min. Sample valve and transfer line were heated at 110 and 120°C respectively. One extraction of the headspace was taken in single puncture mode. Pressure was applied for 0.2 min in order to fill the sample loop, which was then equilibrated for 0.1 min before
injection (3 min) into the GC-MS system. Then GC analysis took place in injection split mode (ratio 10:1). Helium was used as carrier gas with a constant flow of 1.0 ml/min. The column was heated from 40°C (4 min initial time) to 240°C at a rate of 4°C/min and hold at 240°C for 20 min. The mass spectrometer was operated in electron impact mode with source and quadrupole temperatures of respectively 230 and 150°C. Masses were scanned from 20 to 250 Da.

Analysis of aqueous solutions:

One ml of aqueous solution (A1) obtained as described in Example 1 or reconstituted by dissolution of 60 mg of freeze dried powder (A1F) in 1 ml of distilled water, was introduced into a vial for automated headspace sampler. As it is recommended not to heat the samples more than 20°C below boiling temperature to avoid over pressure problems when injecting, samples were preheated in an oven set at 100°C for 2h30 and rapidly cooled down in a bath of ice. The vials were then introduced into the headspace autosampler and reheated at 80°C for 10 min. The temperatures of sample valve and transfer line were respectively 90 and 100°C while all other conditions were identical to those used for the analysis of powders.

The results regarding the aromatic compounds profile is presented in figures 1 to 4.

Figure 1 exhibits Aldehydes generated by heat treatment of A1F powder (■) for 45 min or A1 aqueous solution (□) for 2h30, at 100°C.
Figure 2 exhibits Ketones generated by heat treatment of AlF powder (■) for 45 min or Al aqueous solution (□) for 2h30, at 100°C.

Figure 3 exhibits Furane derivatives generated by heat treatment of AlF powder (■) for 45 min or Al aqueous solution (□) for 2h30, at 100°C.

Figure 4 exhibits Alkyl-pyrazines generated by heat treatment of AlF powder (■) for 45 min or Al aqueous solution (□) for 2h30, at 100°C.

EXAMPLE 8
Application to pizza dough models

In order to evaluate the potential of Al (and AlF) in a product, some application trials on pizza dough were carried out.

Preliminary trials: round table tasting

Two ways of application of the Al aromatising composition were tested: mixed into the dough and as surface coating. The amount of the Al solution added was the same in all samples, corresponding to 1 g of solution per 50 g of fresh dough. In case of mixture with other ingredients of the dough, the Al solution replaced a part of the water involved in the recipe, in order to keep the final moisture content similar for both sample and reference.
The pizza doughs were prepared according to the following recipe:

- Wheat flour: 100 g
- Water: 23 g
- Sunflower oil: 6.8 g
- Salt: 3.0 g
- Baker's yeast: 30 g

All the ingredients were mixed for 210 seconds in a kneading machine. The mixture was then fermented for 25 min in a fermentation cabinet set at 35°C and 85% of relative humidity. The obtained dough was laminated to 5 mm thickness, cut as 10 cm diameter raw pizza crusts and fermented for another 25 min period. Finally the samples were docked and prebaked for 8 min at 220°C before storage at -25°C until the sensory evaluation day.

In addition to Al solution, a preheated Al solution (2 hours at 100°C) was tested in order to check if the baking conditions were sufficient, insufficient, or detrimental to the generation of bread aroma. Two tasting sessions were organized to select the best sample to be evaluated in triangle test versus a reference.

Session 1: Pizza dough, Al mixed into the dough
Session 2: Pizza dough, Al applied as surface coating

In each session, three different pizza crusts were presented: the reference and two spiked samples, one with Al solution and the other one with the preheated Al solution. Immediately after baking (8 min, 200°C), the products were placed under bell-covers fitted with stoppers
in order to trap the aroma in the head-space. The 5-6 panelists were asked to compare the different products, focussing on the smell.

This preliminary round table screening revealed the trend of an increased and pleasant bread crust aroma in most of the spiked samples. The sample treated with preheated Al solution was also considered as pleasant and exhibiting a pleasant bread type aroma, but however slightly weaker than the other samples treated with non-preheated solution.

For all these reasons, the model pizza dough with non preheated Al solution mixed into the dough was selected for further trials.

**Triangle tests**

Triangle tests were performed to verify the impact of the nature of the aromatising composition (liquid or powder). Equivalent amounts of Al or AlF (1.0 g of solution or 60 mg of powder per 50 g of fresh dough), were mixed with other ingredients of the recipe of the pizza dough. The pizza crust were placed under bell-covers immediately after baking. 12 panelists were asked to identify which pizza was different out of the three presented following two schemes: 1 reference, 2 spiked samples or 2 references, 1 spiked sample.

The results were exactly the same for both Al solution and AlF powder. Nine panelists among 12 gave correct answers, meaning that spiked samples were significantly different from the reference with a confidence level of 95%. The panelists were also asked to make free comments to qualify the pizza breads. The main attributes describing the spiked
samples were biscuit, bread crust, cracker and buttery, when yeasty and flour-like were attributed to the reference.

As a general conclusion about these application trials, Al or AlF brought a noticeable pleasant and regularly identified bread crust aroma. Moreover, no off-note was detected throughout the different tasting sessions.
CLAIMS

1. Process for the preparation of an aromatising composition, which comprises the bioconversion of at least two amino compounds selected from the group consisting of amino acids and peptides and at least one reducing sugar in the presence of a micro-organism selected in the group consisting of yeasts.

2. Process according to claim 1, which comprises a separation step to remove the micro-organism and obtaining a supernatant comprising the aromatising composition.

3. Process according to claims 1 and 2, wherein the amino acids are selected in the group consisting of arginine, citrulline, glutamine, ornithine and proline.

4. Process according to claims 1 and 2, wherein the peptides are selected in the group of dipeptides and tripeptides.

5. Process according to claims 1 and 2, wherein the reducing sugar is selected in the group consisting of C5 or C6 monosaccharide.

6. Process according to claim 5, wherein the sugar is selected in the group consisting of fructose, glucose and rhamnose.

7. Process according to claims 1, wherein the micro-organism is selected in the groups consisting of Saccharomyces cerevisiae, Saccharomyces bayanus, Candida versatilis, Debaryomyces hansenii.

8. Process according to claims 1 and 2, wherein the molar ratio between the amino compounds and the reducing sugar is from 1:1 to 1:10.
9. Process according to claim 1, wherein the bioconversion is carried out during 2 to 48 hours, at a pH of 5 to 8 and at a temperature of from 20 to 50°C.

10. Process according to claim 2, wherein the supernatant is dried into a powder.

11. Process for generating baked aroma comprising the step of heating the composition obtainable by a process according to any proceeding claims.

12. Process according to claim 11, wherein the heating treatment is carried out at a temperature of from 90 to 200°C during 5 to 360 minutes.

13. Method for making a bakery product having an improved aroma comprising the steps of
   - mixing flour, water, yeast and an aromatising composition obtainable by a process according to claims 1 to 10,
     - kneading all ingredients in order to obtain a dough,
     - ferment the dough and
     - bake the dough.

14. Method for making a bakery product having an improved aroma comprising the steps of
   - mixing at least flour, water and an aromatising composition obtainable by a process according to claims 1 to 10,
     - kneading all ingredient in order to obtain a dough and
     - bake the dough.

15. Use of an aromatising composition obtainable by a process according to claims 1 to 10 in fermented or non
fermented doughs in order to impart to such products a typical baked aroma.
**INTERNATIONAL SEARCH REPORT**

A. **CLASSIFICATION OF SUBJECT MATTER**

IPC 7 A23L1/23 A21D2/00

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)

IPC 7 A23L A21D

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 practical, search terms used)

EPO-Internal, BIOSIS, FSTA

C. **DOCUMENTS CONSIDERED TO BE RELEVANT**

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<th>Category *</th>
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<tr>
<td></td>
<td>US 3 466 176 A (BUNDUS ROBERT H ET AL) 9 September 1969 (1969-09-09) claims; example 8</td>
<td>1.3-7, 11-15</td>
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<td>EP 0 951 841 A (NESTLE SA) 27 October 1999 (1999-10-27) paragraph ‘0006’ - paragraph ‘0008’; example 1 paragraph ‘0010’ - paragraph ‘0016’; claims</td>
<td>1.2,4-7, 9-15</td>
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<td>US 5 108 766 A (GELINAS PIERRE ET AL) 28 April 1992 (1992-04-28) claims; example 1</td>
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* Special categories of cited documents:

*A* document defining the general state of the art which is not considered to be of particular relevance

*B* earlier document but published on or after the International filing date

*L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

*O* document referring to an oral disclosure, use, exhibition or other means

*P* document published prior to the international filing date but later than the priority date claimed

* †* 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

*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 to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.

* &* document member of the same patent family

Date of the actual completion of the International search 13 November 2002

Date of mailing of the international search report 20/11/2002

Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL-2280 H-V P fulfilw. - Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax (+31-70) 340-3016

Authorized officer Lepretre, F
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