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(54) METHOD FOR OPERATING A SYSTEM FOR THE INTEGRATED AND AUTOMATED ANALYSIS OF DNA OR PROTEIN

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- (56) **References Cited**

U.S. PATENT DOCUMENTS

5,096,669 A	3/1992	Lauks et al.	
5,755,942 A	5/1998	Zanzucchi et al.	
	(Continued)		

FOREIGN PATENT DOCUMENTS

DE	4438785 A1	5/1996
DE	19648695 A1	6/1997
	(Conti	nued)

OTHER PUBLICATIONS

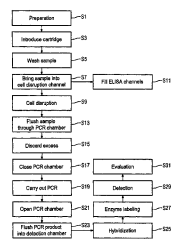
Buch et al. Anal Chem. 2004. 76:874-881.* (Continued)

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(57) **ABSTRACT**

A method for the integrated and automated analysis of DNA or protein, including a single-use cartridge, with an analysis device having a control device, and devices for capturing and processing signals, the control device carrying out a completely automatic process and evaluation of molecular diagnostic analysis via single-use cartridges (Lab-on-a-Chip). Controlling of an analysis process, which occurs in the cartridge, involve the subsequent displacement and thermostatisation of liquids with a first device, and with a second device the signals which are obtained during the analysis are processed. The first and the second devices are synchronized in such a manner that the analysis process of the sample can be carried out in a totally integrated manner thus producing an immediate result.

34 Claims, 9 Drawing Sheets



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(56)**References** Cited

U.S. PATENT DOCUMENTS

5,764,356	Α	6/1998	Iwase et al.
6,054,277	Α	4/2000	Furcht et al.
6,300,141	B1	10/2001	Segal et al.
6,458,325	B1	10/2002	Roscher et al.
6,780,300	B1	8/2004	Alberto
6,814,846	B1	11/2004	Berndt
7,060,488	B2	6/2006	Duong et al.
7,122,153	B2	10/2006	Но
2001/0012612	A1	8/2001	Petersen et al.
2002/0019060	A1 $*$	2/2002	Petersen B01L 3/502
			436/514
2002/0042125	A1*	4/2002	Petersen B01L 3/502715
			435/287.2
2002/0142482	A1	10/2002	Wu et al.
2002/0179444	A1	12/2002	Lauks
2003/0073229	A1	4/2003	Greenstein et al.
2003/0148530	A1	8/2003	Lauks
2003/0199910	Al	10/2003	Boecker et al.
2003/0207326	A1	11/2003	Su et al.
2004/0029203	A1*	2/2004	Gumbrecht B01J 19/0046
200 0 0023200		2,200.	435/7.92
2004/0086870	A1	5/2004	Tyvoll et al.
2004/0115094	Al	6/2004	Gumbrecht et al.
2004/0189311	Al	9/2004	Glezer et al.

2004/0223874 A1*	11/2004	Numajiri	B01L 3/502715
			422/400

2004/0224339 A1	11/2004	Numajiri et al.
2005/0031490 A1	2/2005	Gumbrecht et al.
2009/0130658 A1	5/2009	Barlag et al.
2009/0136922 A1	5/2009	Barlag et al.

FOREIGN PATENT DOCUMENTS

DE	19648441 A1	5/1998
DE	19928410 A1	1/2001
DE	10258840 A1	8/2003
DE	10352887 A1	4/2005
EP	0992287 A2	4/2000
EP	1203959 A1	5/2002
EP	1418243 A2	5/2004
EP	1473085 A1	11/2004
WO	1998022816 A1	5/1998
WO	1999010735 A1	3/1999
WO	1999044740 A1	9/1999
WO	1999046045 A1	9/1999
WO	2000052457 A1	9/2000
WO	2001047638 A2	7/2001
WO	2002072262 A1	9/2002
WO	2002073153 A2	9/2002
WO	2002097398 A2	12/2002
WO	2004062804 A1	7/2004
WO	2006042838 A1	4/2006

OTHER PUBLICATIONS

Tisone. IVD Technology Magazine. Originally published May 1998. Retrieved on Sep. 28, 2016 from the internet: http://www. bioon.com/biochip/general/011-2.html.*

James A. Higgins, et al., A Handheld Real Time Thermal Cycler for Bacterial Pathogen Detection, Elsevier Science B.V., Received July. 18, 2002; Received in Revised Form Sep. 26, 2002; Accepted Oct. 24, 2002; pp. 1115-1123.

P.-A. Auroux, et al., Miniaturised Nucleic Acid Analysis, The Journal Is the Royal Society of 2004, Lab Chemistry Chip, 2004, 4, pp. 534-546.

Robin Hui Liu, et al., Self-Contained, Fully Integrated Biochip for Sample Prpearation, Polymerase Chain Reaction Amplification, and DNA Microarray Detection, Analytical Chemistry, vol. 76, No. 7, Apr. 1, 2004, pp. 1824-1831.

Eric T. Lagally, et al., Fully Integrated PCR-Capillary Electrophoresis Microsystem for DNA Analysis, This Journal is the Royal Society of Chemistry 2001, Lab on a Chip, 2001, 1, pp. 102-107. W.J. Benett, et al., Handheld Advanced Nucleic Acid Analyzer, Biochemical and Biomolecular Sensing, Robert A. Liberman, Editor, Proceedings of Spie, vol. 4200 (2000), pp. 55-63.

Magnetic Shielding (retreived on Feb. 11, 2014 from the Internet: http://www.magneticshielding.com/magnetic-shielding-faq.html).

MuMetal (retreived on Feb. 11, 2014 from the internet: http://metals. about.com/library/bldef-Mu-Metal.htm).

Nickel Alloys.net (retreived on Feb. 11, 2014 from the internet: http://www.nickel-alloys.net/nickelalloys.html).

* cited by examiner

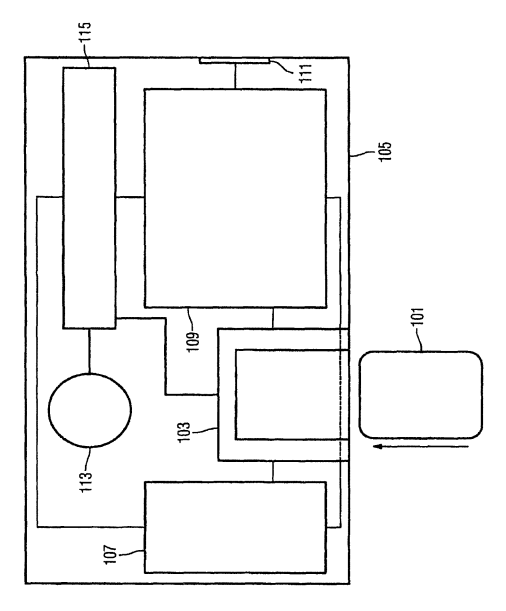
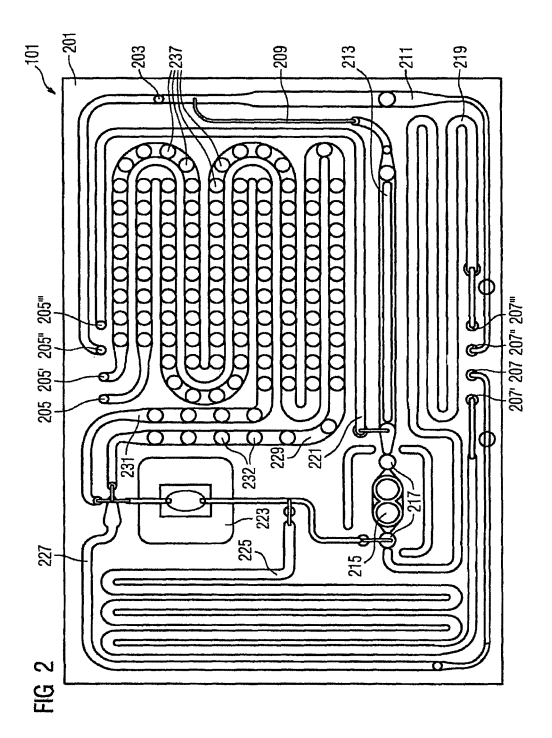
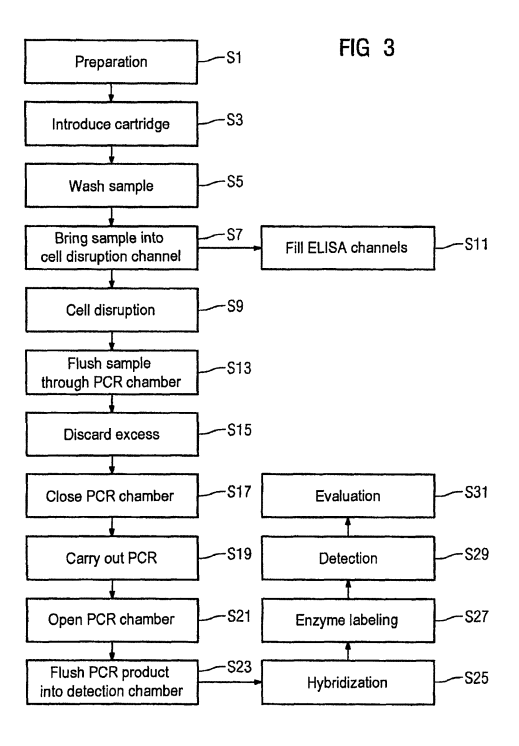
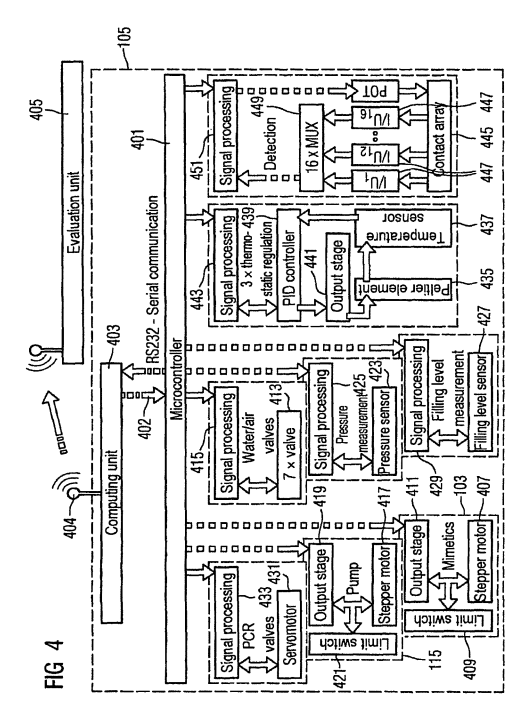
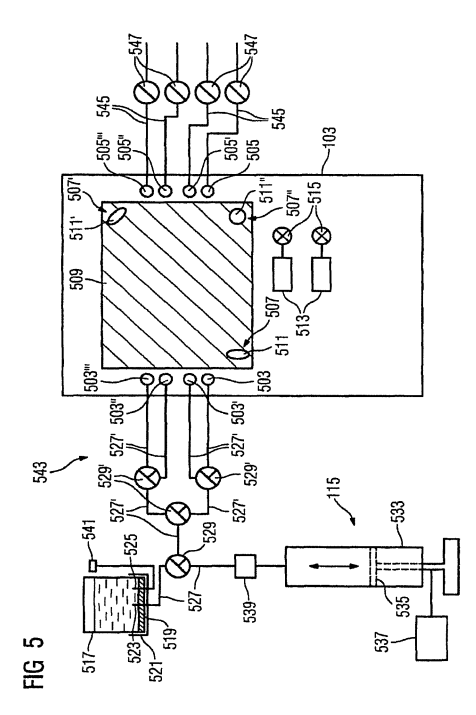


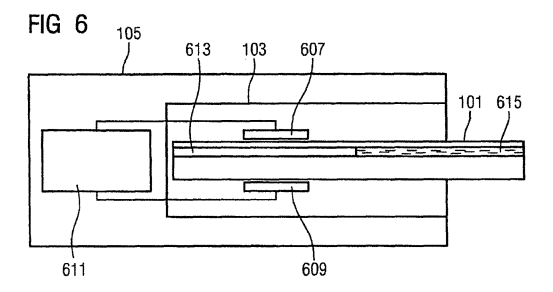
FIG 1

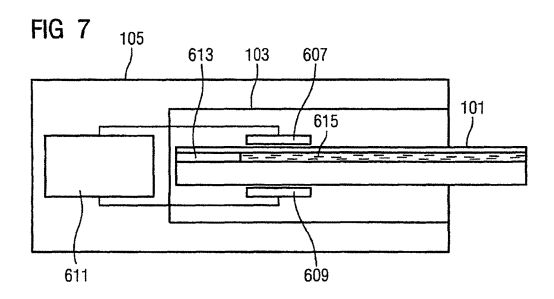


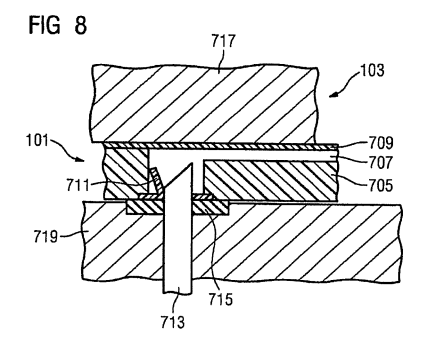


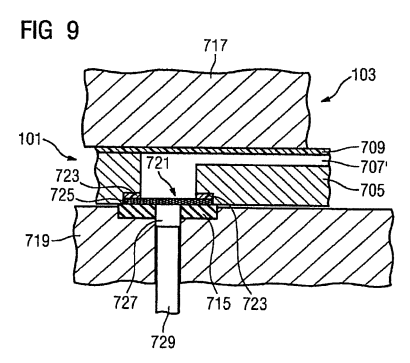


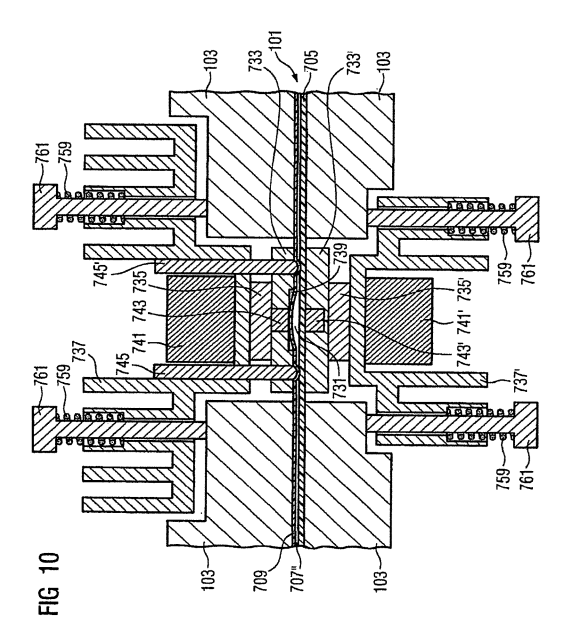


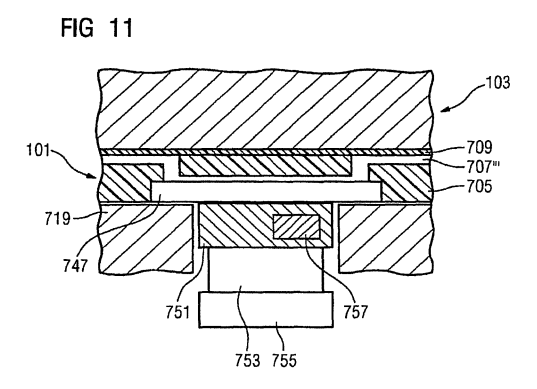












METHOD FOR OPERATING A SYSTEM FOR THE INTEGRATED AND AUTOMATED ANALYSIS OF DNA OR PROTEIN

CROSS REFERENCE TO RELATED APPLICATION

This application is a division of commonly owned, copending U.S. patent application Ser. No. 11/920,978, filed Jul. 23, 2009, which is a § 371 of PCT/EP2006/062499 filed May 22, 2006 and claims 35 USC § 119 priority of German Patent Application DE 10 2008 059 536.7 filed Dec. 17, 2005.

BACKGROUND OF THE INVENTION

Field of the Invention

At least one embodiment generally relates to a system for the integrated and automated analysis of DNA or protein, for $_{20}$ example one comprising a single-use cartridge (card), an evaluation device and a device for signal recording and processing. In particular, at least one embodiment of the invention generally relates to a control device as part of the evaluation device for the fully automatic processing and 25 evaluation of molecular diagnostic analyses with disposable cartridges ("lab-on-a-chip") and also known as a biochip; see, commonly owned U.S. Patent Application Publication 2007/01411553). In addition, at least one embodiment of the invention also generally relates to a method for operating the 30 type of system.

Description of Related Art

The significance of decentralized analysis devices ("point of care", "near patient testing") in molecular diagnostics or general medical technology is increasing. In addition, fast 35 tests are becoming more and more important in the areas of foodstuff monitoring and environmental protection.

The general literature, but also especially the patent literature includes a large number of publications on the aforementioned topic: International Patent Application Pub- 40 lications WO 02/072 262 A1 and WO 02/073 153 A1 (which corresponds to U.S. Patent Application Publication 2005/ 0031490) disclose systems for application in biomedical technology in which samples taken in a decentralized manner are evaluated in a decentralized manner by means of an 45 optionally mobile read-out device. The samples, in particular, are in this case collected in a single-use cartridge formed like a check card ("card").

Such a cartridge for application in the DNA analysis of whole blood samples is disclosed in detail in International 50 Patent Application Publication WO 2006/042 838 A1 and corresponding U.S. Pat. No. 7,851,227, not previously published. In this case, the following features of the cartridge are regarded as essential for automatically carrying out all the measures required for process analysis from individual 55 partial processes in the cartridge:

the cartridge contains a system of microchannels and/or microcavities for a microfluidic process technology,

- the microchannels or the microcavities have predetermined geometrical structures for receiving reagents, 60 wherein
- the reagents are stored in a storage-stable form at specific locations in the microchannels or the microcavities of the cartridge.
- means are present for providing the dry-stored reagents 65 for the respective partial process in a suitable form, in particular as a liquid reagent.

Reference is expressly made to the disclosure of these features in the above earlier application.

Furthermore, U.S. Patent Application Publications 2002/ 0179444 A1 and 2003/0148530 A1 disclose individual solu-

tions for a blood diagnosis method and the associated signal processing. Specific problem solutions for the realization of the system of the type mentioned in the introduction comprising the previously described card and associated suitable means for evaluation are described in earlier applications in 10the name of the applicant that have not yet been published.

SUMMARY OF THE INVENTION

At least one embodiment of the invention is directed to a 15 system which meets practical requirements in particular with regard to the evaluation device. At least one embodiment is directed to an associated control method for evaluation in the case of a system comprising cartridge and evaluation device. In the case of at least one embodiment, the following requirements are made of the evaluation device:

low price

small structural design

simple operating control ("one button" operation)

rapid performance of the analysis

integrated microfluidics

safety against contamination, i.e., avoidance of instances of soiling and infections-high operational reliability

capability of communication with data processing systems.

No practically suitable solution has been disclosed hitherto for the fully automatic nucleic acid analysis from taking the sample (e.g., blood) to displaying the result, in particular including cell disruption, PCR and detection. Since the equipment currently proposed for biochemical analysis does not yet meet all the practical requirements, at least one embodiment of the invention improves the system so as to allow a maximally automated and reproducible evaluation. In particular, the intention here is to ensure that the result is displayed directly on a display or the data are forwarded into a computer system. The intention is to specify, in at least one embodiment, a suitable operating method for this purpose.

The operating method of at least one embodiment for carrying out an evaluation of a sample in a single-use cartridge with the aid of the control device in a system. Especially the realizations for carrying out a DNA analysis, and also in particular an SNP analysis, or alternatively a protein analysis, are contained in further embodiments of the method.

The basis of at least one embodiment of the invention, therefore, is an improved concept by comparison with the prior art: a device construction realized in accordance with this concept is designed according to at least one embodiment of the invention for the fully automatic processing and evaluation of molecular diagnostic analyses with disposable cartridges (lab-on-a-chip), in particular for the control and read-out of analysis processes within cartridges in the form of modified check cards.

In the case of at least one embodiment of the invention, the control device has the following features, in particular:

- a cartridge receptacle for receiving, locking, unlocking and releasing the cartridge,
- a device for correctly positioning the cartridge in the reader.
- sensors for identifying the cartridge, in particular a correct cartridge, e.g., limit switches, barcode,
- a water reservoir for diluting the sample, dissolving dry reagents of the cartridge, carrying out flushing opera-

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tions, wherein fluidic contact can be made with the water reservoir by way of a septum being pierced by a pointed hollow needle,

- a sterile filter ventilation valve for ventilating the water reservoir when water is drawn,
- a power supply unit,
- a pump for conveying extremely small quantities of liquid, for example in the microliters range,
- means for distributing water via valves, e.g., water-carrying solenoid valves,
- a pressure sensor in the water feed for monitoring fluidic operations,
- at least one interface for feeding water into the cartridge,
- sensors for identifying the filling of throughflow channels on the cartridge, e.g., capacitive sensors with electrodes 15 above and below the cartridge,
- a device for closing off and opening venting openings of the cartridge, e.g., air-carrying solenoid valves,
- a device for closing off and opening chambers, in particular the PCR chamber, on the cartridge, e.g., plung- 20 ers pressing onto a film,
- a combination comprising a magnetic bead collector and a thermostatic regulation device for capturing magnetic beads and for carrying out thermocycling processes for the PCR,
- a sandwich arrangement of two abovementioned devices for rapid thermocycling,
- a device for ensuring optimum heat transfer between the thermostatic regulation device and cartridge, e.g., by way of a contact pressure defined by way of springs,
- a device for making electrical contact with an electrical detection module, e.g., contact pins,
- an integrated thermostatic regulation for the detection module,
- control, measuring and regulating electronics for all the 35 functions.
- a microcontroller for control and data acquisition, and
- a display for representing the analysis results.

In the associated method for operating the control and read-out device for the evaluation of a cartridge filled with 40 a sample, the following method steps are carried out specifically for a DNA analysis:

- the cartridge receptacle is opened and the pump draws water from the reservoir given a suitable valve position, the control device indicates readiness for operation,
- a user inserts the cartridge with a sample, e.g., blood, the analysis is started by a "one button" action or auto-
- matically, e.g., by way of a cartridge limit switch, given a suitable valve position, water is pumped into the
- cartridge via a water port in order to wash out excess 50 sample material,
- the venting valves are actuated in a suitable manner in order to allow or prevent the filling of cartridge channels by way of permitting or avoiding venting,
- as a result of the inflow of water, the sample is diluted and 55 pumped into the cell disruption channel, wherein dry reagents are dissolved,
- a residence time ensues for cell disruption, wherein liberated DNA binds to magnetic beads,
- nearly at the same time two ELISA reagent channels are 60 filled, wherein dry ELISA reagents (enzyme label and substrate) are dissolved,
- the content of the cell disruption channel is flushed by further pumping of water through the PCR chamber, wherein the magnetic beads with DNA are held back by 65 a magnetic field, the remainder is flushed into a waste channel,

the valves of the PCR chamber are closed off,

- the thermocycling is started, wherein dry PCR reagents are dissolved,
- a predetermined number of thermocycles are carried out at defined temperatures (PCR),
- the valves of the PCR chamber are opened after conclusion of the PCR,
- given a suitable valve position, water is pumped into the cartridge via a further water port and the PCR product is flushed from the PCR chamber into a detection chamber,
- the detection chamber is thermostatically regulated, wherein hybridization of the DNA with catcher molecules takes place,
- air possibly present between the PCR product and the dissolved ELISA reagents is pumped into a waste channel by suitable setting of water port and venting valves,
- the dissolved enzyme label is pumped via the detection chamber into a further waste channel by suitable setting of water port and venting valves,
- the dissolved substrate is pumped via the detection chamber into a further waste channel by suitable setting of water port and venting valves,
- the electrical detection is started,
- e.g., an SNP analysis is carried out by recording melting curves.

As an alternative to the above method sequence, a protein analysis is also possible, wherein no cell disruption and no PCR are necessary. In this case, the proteins sought are isolated from the sample by way of magnetic beads and detected in the detection chamber. Haptens can be detected analogously.

An analysis process integrated throughout is carried out, then, by the method according to an embodiment of the invention. This is understood to be a fully automated method sequence without manual intervention. After the introduction of the cartridge with the sample into the control device, the sample treatment and analysis process proceeds automatically until the actual detection of the substances sought. This has the advantage, in particular, of high reproducibility and a low error rate.

What is particularly advantageous in the system according to at least one embodiment of the invention is that the control device can be embodied in compact fashion and has both internal fluidic interfaces and electrical interfaces. Interfaces for linking to external computer systems are additionally present. These may be serial and/or wireless interfaces, for example. In an example embodiment, the control device comprises a connection possibility for a commercially available PDA. This enables a display to be integrated into the control device in a simple manner. In a particularly preferred example embodiment, a display and operating unit is fixedly integrated into the control device.

With the system according to at least one embodiment of the invention, a considerable advance is achieved compared with the previously proposed devices. It is now possible, for example, for the measured data to be directly displayed or forwarded for evaluation.

Further details and advantages of the application are explained on the basis of the example embodiment described below in association with the accompanying drawings, in which:

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 shows a schematic overview diagram of an analysis device,

FIG. **2** shows a schematic overview diagram of a cartridge,

FIG. 3 shows a schematic flowchart of an analysis method,

FIG. **4** shows a schematic overview diagram of the ⁵ functional units of the control device,

FIG. **5** shows a schematic overview diagram of the fluidic components of the control device,

FIG. 6 and FIG. 7 show schematic illustrations of a measuring device for a filling level,

FIG. **8** shows a schematic illustration of a fluidic interface between the cartridge and the control device,

FIG. 9 shows a schematic illustration of a venting interface between the cartridge and the control device,

FIG. **10** shows a schematic illustration of a PCR chamber ¹⁵ of the cartridge, and

FIG. **11** shows a schematic illustration of a detection chamber of the cartridge.

DETAILED DESCRIPTION OF THE INVENTION

An analysis device is described below as one example embodiment of the invention. The analysis device is divided into a control device and a cartridge. Whereas the cartridge 25 is, for example, present as a disposable article on which the actual analysis process proceeds, the control device is reusable and serves for controlling the analysis process. The cartridge contains only passive components; the entire control proceeds from the control device. 30

FIG. 1 shows the construction of the analysis device. It includes a cartridge 101, which can be inserted into a cartridge receptacle 103 of a control device 105. The cartridge receptacle 103 is connected to a power supply unit 107, which supplies the receptacle with power. For the 35 transmission of data of the analysis process, the cartridge receptacle 103 is additionally connected to an electronic unit 109. The data can be output from the electronic unit 109 via an interface 111.

The interface **111** is formed as an RS232 interface, for 40 example. The electronic unit **109** is connected to the power supply unit **107** for its power supply.

In order to carry out the analysis process, water is required in the cartridge. This water is stored in a supply container **113**. The supply container **113** is a commercially available 45 bottle containing sterile, distilled water, for example, which is closed off by a septum. A corresponding receptacle for the bottle in the control device is provided with a cannula which pierces the septum when the bottle is inserted. As a result, the water becomes accessible and it can be drawn from the 50 bottle via a connected pump **115**. The pump **115** can take up a defined quantity of water and pump it into the cartridge receptacle **103** via a connection. The water passes into the cartridge **101** via the cartridge receptacle **103** and is available in the analysis process. The pump **115** is connected to 55 the power supply unit **107** for its power supply.

FIG. **2** schematically illustrates the cartridge **101**, which has already been disclosed in detail in International Patent Application Publication WO 2006/042 838 A1 and corresponding U.S. Pat. No. 7,851,227, not previously published, 60 which disclosure is also subject matter of the present patent application, wherein the entire contents thereof is hereby incorporated herein by reference.

The cartridge **101** is present in the form of a plastic card **201** of the size of a check card and comprises a number of 65 channels. Via a sample port **203**, a sample can be pipetted into the cartridge **101**. Water can be introduced into the

cartridge 101 via four water ports 205, 205', 205" and 205"". The cartridge 101 is additionally provided with four venting ports 207, 207', 207" and 207", through which air can escape when the channels are filled. A metering section 209 comprises a volume of 1 1 and services for tapping of this quantity from the sample that has been pipetted in. A sample waste channel 211 serves for receiving the rest of the sample. The venting port 207" lies at the end of the sample waste channel 211. The metering section 209 is adjoined by a cell disruption channel 213, in which reagents for the cell disruption are stored in dry form. It is only as a result of the introduction of the sample in diluted form that the reagents are dissolved and cell disruption takes place. As a result of the cell disruption, DNA contained in the cells is liberated, said DNA binding to magnetic beads likewise present in the cell disruption channel 213.

The cell disruption channel 213 is connected to a PCR chamber 215, in which a PCR can be carried out. The PCR chamber 215 can be closed off by way of two valves 217. 20 The required reagents are situated in the PCR chamber 215 likewise in dry form. Arranged downstream of the PCR chamber 215 is a PCR waste channel 219 connected to the venting port 207". A water channel 221 connected to the water port 205" leads in at one end of the cell disruption channel 213. The PCR chamber 215 is connected to a detection chamber 223, in which the DNA can be detected by means of a sensor array. The detection chamber 223 is connected to two waste channels 225 and 227 ending in the ventilation ports 207' and 207, respectively. In addition, the detection chamber 223 is connected to two ELISA reagent channels 229 and 231 (ELISA: enzyme linked immuno sorbent assay), which, for their part, are connected to the water ports 205' and 205, respectively. ELISA reagents 232 are stored in dry form in the ELISA reagent channels 229 and 231, and after being dissolved in water can be flushed into the detection chamber 223.

FIG. 3 shows a schematic flowchart of an analysis process. In a first method step S1, the cartridge is prepared. This involves pipetting the sample into the cartridge and possibly providing water in the control device. As it is pipetted in, the sample accumulates in the sample waste channel 211 of the cartridge (referring to FIG. 2), the metering section 209 likewise filling up in the process. In a second method step S3, the cartridge is introduced into the control device. As a result of the introduction of the cartridge, valves situated in the read-out device are connected to the water ports 205 to 205''' and the venting ports 207 to 207''', such that water can be introduced into the cartridge and air can escape. Moreover, a switch in the read-out device is actuated, whereupon the analysis process starts automatically.

In this case it is ensured by way of functional elements known per se that the cartridge is in the correct position and has the correct orientation. Moreover, the fact of whether the cartridge is one which is suitable for the control device is identified. At the beginning of the analysis process within the cartridge 101, the water channel 221 is filled with water, the water channel 221 likewise being vented via the PCR waste channel 219. This simplifies the method sequence since the air present in the water channel 221 could not escape in the further sequence.

In a third method step S5, water is pumped into the water port 205", such that that portion of the sample which is situated in the metering section 209 is isolated. The venting port 207", which is otherwise closed off by the valve, is opened for this purpose. As soon as the sample waste channel 211 has been filled, the venting port 207" is closed off again in a fourth method step S7. The filling level of the

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sample waste channel **211** is measured by way of filling level sensors arranged in the control device. The functioning is explained with reference to FIGS. **6** and **7**.

As a result of the venting port **207**["] being closed off, pressure starts to build up as water continues to be pumped 5 in, such that that portion of the sample which is situated in the metering section **209** is flushed into the cell disruption channel **213**. The valve assigned to the venting port **207**" in the control device is opened for this purpose. The sample is diluted with water. As a result of the introduction of the 10 diluted sample, the dry reagents in the cell disruption channel **213** are dissolved whereupon cell disruption takes place in a fifth method step **S9**. DNA situated in the cells of the sample is liberated in this case. The liberated DNA binds to magnetic beads stored in the cell disruption channel **213**. 15

At the same time as the cell disruption, in a sixth method step S11, the ELISA reagent channels 229 and 231 are filled with water. The valves assigned to the venting ports 207' and 207 in the control device are opened for this purpose. The filling level of the ELISA reagent channels 229 and 231 is 20 measured by filling level sensors arranged in the control device.

After the cell disruption has been concluded, in a seventh method step S13, the diluted sample with the magnetic bead-DNA complexes is flushed through the PCR chamber 25 215. For this purpose, as already previously, water is pumped through the water port 205" and the venting port 207" is opened. In the control device, there is a magnet arranged above the PCR chamber 215, the magnetic bead-DNA complexes being held back in the PCR chamber 215 30 by said magnet. The remainder of the diluted sample is flushed into the PCR waste channel 219. In an eighth method step S15, water is again pumped through the PCR chamber 215, thereby cleaning the magnetic bead-DNA complexes.

In a ninth method step S17, the valves 217 of the PCR 35 chamber 215 are closed. In a tenth method step S19, the DNA of the magnetic bead-DNA complexes is replicated by a PCR. For this purpose, corresponding reagents are situated in the PCR chamber 215 in dry form, said reagents being dissolved by the addition of water. In order to manipulate the 40 temperature in the PCR chamber 215, two Peltier units are situated in the control device. After the conclusion of the PCR, in an eleventh method step S21, the valves 217 of the PCR chamber 215 are opened. In a twelfth method step S23, water is pumped into the channel 221 via the water port 45 205", such that the water previously situated in the channel 221 is flushed into the PCR chamber 215. The channel 221 contains dry stored salts which are thus added to the PCR product, that is to say the replicated DNA. At the same time, the replicated DNA is flushed from the PCR chamber 215 50 into the detection chamber 223. In this case, the venting ports 207" and 207' are closed, and the venting port 207 is open.

In a thirteenth method step S25, a hybridization between the PCR product and capture molecules arranged in the detection chamber 223 takes place in the detection chamber 223. In order to support the hybridization, the temperature in the detection chamber 215 is set in a defined manner by way of a Peltier element arranged in the control device. In a fourteenth method step S27, a label enzyme (e.g., streptavidin alkaline phosphatase, streptavidin esterase or oligonucleotide-coupled esterase) linked to a coupling molecule is pumped from the ELISA reagent channel 231 through the detection chamber. For this purpose, water is pumped through the water port 205 and the venting port 207' is by the label enzyme. In a fifteenth method step S29, a

dissolved enzyme substrate is pumped from the ELISA reagent channel **229** through the detection chamber **215**. Both enzyme label and enzyme substrate channel are vented before the flooding of the detection chamber **231** via the channel **227** in order to prevent air from being entrained into the detection chamber **223**. By virtue of the substrate, a so-called redox cycling process occurs in the detection chamber, and the bound DNA can be detected by said process. Evaluation of the measurement data correspondingly takes place in a sixteenth method step **S31**.

FIG. 4 illustrates the central components of the control device 105 in a block diagram. A microcontroller 401 serves as central controller of the entire analysis process, said microcontroller being connected to a computing unit 403 via a serial RS232 interface 402. Said computing unit 403 is a commercially available PDA, for example, and serves for recording measurement data from the analysis process. The computing unit 403 is equipped with a radio module 404, via which the measurement data can be transmitted to an evaluation unit 405. The radio module 404 communicates with the evaluation server 405 for example by infrared, Bluetooth or WLAN.

The microcontroller **401** controls all the components of the control device **105** which are involved in the analysis process. The cartridge receptacle **103** comprises a stepper motor **407**, which moves a part of the cartridge receptacle **103** upon insertion of the cartridge, such that the cartridge is correctly received. The Peltier elements are brought into position in this case. A limit switch **409** interrupts the process when the cartridge has been completely introduced. As a result of the introduction process, the water ports and the venting ports are connected to corresponding channels in the control device. Valves **413** controlled by the microcontroller **401** are arranged in the channels. A signal processing unit **415** serves for communicating the control signals. The arrangement and functioning of the channels and valves are explained with reference to FIG. **5**.

For the movement of the water in the channels, the pump **115** is controlled by the microcontroller **401**. Its piston is moved by a stepper motor **417**, which is in turn controlled by means of an output stage **419** and a limit switch **421**. The pressure in the pump **115** is measured by means of a pressure sensor **423** and read out by the microcontroller **401** via a signal processing unit **425**.

The movement of the water in the cartridge is monitored by means of filling level sensors **427**, which are read by the microcontroller **401** via a signal processing unit **429**.

The valves of the PCR chamber are controlled via a servomotor **431** and a signal processing unit **433**. Three Peltier elements **435** and temperature sensors **437**, which are controlled and read via a PID controller **439** and an output stage **441**, serve for the thermostatic regulation of the PCR chamber and the detection chamber. The PID controller **439** is controlled by the microcontroller **401** via a signal processing unit **443**.

The signals generated by the redox cycling on the sensor array are measured via a contact-making module **445** and forwarded to a signal processing unit **451** via read-out circuits **447** and a multiplexer **449**. The signal processing unit **451** is connected to the microcontroller. The necessary control voltages, such as e.g., a reference voltage and a counter-voltage of a potentiostat, are fed to the sensor array via the contact-making module **445**. The sensor array is embodied as a CMOS sensor chip, for example. In this case, the contact-making module **445** advantageously has the arrangement as in a commercially available smart card reader.

FIG. 5 illustrates a schematic overview diagram of the fluidic components of the control device. The cartridge receptacle 103 includes four water ports 503, 503', 503" and 503", which are connected to the water ports of the cartridge when the latter has been inserted. Four venting ports 505, 505', 505" and 505" are likewise provided, which are connected to the corresponding venting ports of the cartridge. The cartridge receptacle 103 furthermore includes three filling level sensors 507, 507' and 507", the functioning of which is explained with reference to FIGS. 6 and 7. They operate capacitively and comprise two capacitor plates. In order to ensure a simple construction of the cartridge receptacle 103, one capacitor plate 509 is formed in large-area fashion and is used jointly for all the filling level sensors 507, 507' and 507". The other capacitor plates 511, 511' and 511" are correspondingly embodied separately.

The filling level sensors 507 and 507' are designed in each case for monitoring two channels of the cartridge and are arranged at corresponding locations of the cartridge recep- 20 tacle 103. The filling level sensor 507 serves for monitoring the two ELISA reagent channels 229 and 231, while the filling level sensor 507' monitors the sample waste channel 211 and the PCR waste channel 219. The filling level sensor 507" serves for monitoring the waste channel 227.

Two motors 503 for controlling two actuators 515 are additionally arranged in the cartridge receptacle 103. The actuators 515 serve for opening and closing the valves of the PCR chamber on the cartridge.

The water required for the analysis process comes from a 30 commercially available bottle 517, which is closed off by a septum 519. The bottle 517 is held by a receptacle 521 of the control device, two cannulas 523 and 525 being arranged in said receptacle. When the bottle 517 is introduced into the receptacle 521, the septum 519 is pierced by the two 35 cannulas 523 and 525 and the water in the bottle 517 is made accessible. The cannula 523 is connected to a pump 115 via a line 527 and a three-way valve 529. By means of the pump 115, a volume of water that can be set in a defined manner can be drawn from the bottle 517. The pump 115 comprises 40 a hollow cylinder 533 with a movable piston 535. The piston 535 can be moved in a defined manner by means of a stepper motor 537. A pressure sensor 539 is arranged in the line 527, and the pressure in the line 527 can be determined by way of the pressure sensor. 45

The cannula 525 serves for ventilating the bottle, since a negative pressure would otherwise arise when water is drawn. The cannula is connected to a sterile filter 541, via which air can pass into the bottle.

The three-way valve 529 can be connected to the four 50 water ports 503, 503', 503" and 503" via a line system 543. A plurality of lines 527' and three further three-way valves 529', by which the three-way valve 529 can be selectively connected to each of the water ports 503, 503', 503" and 503", are provided for this purpose. Through suitable 55 switching of the three-way valves 529', the water can thus be pumped from the hollow cylinder 533 into one of the water ports 503, 503', 503" or 503". The pressure occurring in the line system 543 and the line 527 is detected by the pressure sensor 539 in this case. 60

The air present in the channels of the cartridge before the start of the analysis process must be able to pass out of the cartridge as the latter is filled with water. The four ventilating ports 505, 505', 505" and 505" are provided for this purpose in the cartridge receptacle. They are connected via 65 lines 545 to two-way valves 547, which can be optionally closed or open. When water is introduced into one of the

channels, the corresponding one of the valves 547 is opened, such that air can escape from the cartridge.

FIGS. 6 and 7 show the functioning of the filling level sensors. FIG. 6 illustrates the control device 105 in a sectional view. A cartridge 101 has been introduced into the cartridge receptacle 103. In the cartridge receptacle 103, two capacitor plates 607 and 609 are arranged above and below the cartridge 101. The capacitor plates 607 and 609 are connected to a measuring unit 611, by which the capacitance of the capacitor formed by the capacitor plates 607 and 609 can be determined.

In FIG. 6, a channel 613 shaped within the cartridge 101 is partly filled with water 615. In the illustration in FIG. 7, the channel 613 is filled to an extent such that there is water 615' between the capacitor plates 607 and 609. As a result, the capacitance of the capacitor is altered in comparison with the state illustrated in FIG. 6. The change is determined by means of the measuring unit 611 and the liquid level in the channel is determined in this way.

FIG. 8 shows a section through a part of the cartridge receptacle 103 with cartridge 101 inserted. The contactmaking between the water ports of the control device 103 and the cartridge 101 is shown. A channel 707 is milled in a plastic body 705 of the cartridge 101. The channel 707 is covered by way of an adhesive film 709, the adhesive film 709 being adhesively bonded onto the plastic body 705. The adhesive film 709 is commercially available and compatible with all the processes that proceed in the cartridge 101. A further adhesive film 711 is fixed to the water port of the cartridge 101. The further adhesive film can be pierced by a hollow needle 713 arranged in the cartridge receptacle 103 and the fluidic contact between the cartridge receptacle 103 and the cartridge 101 can thus be established. A seal 715 that prevents liquids from escaping is arranged in the cartridge receptacle 103. the seal 715 is preferably composed of a biochemically compatible silicone material in order to avoid contamination of the water ports.

Before the cartridge 101 is introduced into the cartridge receptacle 103, an upper part 717 and a lower part 719 of the cartridge receptacle 103 are further away from one another than illustrated in FIG. 8. This enables the cartridge 101 to be introduced in a convenient manner. As soon as the cartridge 101 has been completely introduced, the two parts 717 and 719 are brought together and the cartridge 101 is locked in the cartridge receptacle 103. At the same time, the adhesive film 711 is pierced by the hollow needle 713.

FIG. 9 shows a section through a part of the cartridge receptacle 103 with inserted cartridge 101 at one of the venting ports. In the cartridge 101, analogously to the illustration in FIG. 8, a channel 707' is milled into the plastic body 705, said channel being covered by the adhesive film 709. Around the venting port 721 of the cartridge 101, a depression is milled into the plastic body 705, an adhesive layer 723 being arranged in said depression. A Teflon membrane 725 is fixed on the adhesive layer 723. A seal 715 is arranged in the lower part 719 of the cartridge receptacle 103. A channel 727 is additionally provided, which runs through the seal 715 and is therefore in contact with the Teflon membrane 725. A hose 729 connected to the venting valves of the control device is arranged in the channel 727.

The Teflon membrane 725 ensures that no liquid can escape from the cartridge. This serves to avoid contamination of the control device, which is intended to be able to be used for a multiplicity of analysis processes. The sample introduced into the cartridge 101 generally contains infectious biomaterial which would contaminate the control device upon contact, and the control device would therefore

have to be cleaned or replaced for carrying out further analysis processes. The Teflon membrane **725** is permeable to vapor and gas, but impermeable to liquids. By way of the pressure sensor **533** of the control device it is ensured that the pressure in the cartridge **101** does not become excessively high and the Teflon membrane **725** or the adhesive film **709** is not damaged or torn.

FIG. 10 illustrates a section through a part of the cartridge receptacle 103 with inserted cartridge 101. Analogously to FIGS. 8 and 9, a channel 707" is milled into the plastic body 705. The channel 707" has a widening serving as a PCR chamber 731. The channel 707" and the PCR chamber 731 are covered by the adhesive film 709.

In order to ensure rapid heat transfer for the thermocycling during the PCR, the PCR chamber **731** is kept as small as possible in a geometrical dimension (approximately 1 mm or volume <20 μ l) and the heat transfers are realized in a "sandwich" thermostatically regulating arrangement such that only small liquid layers (a few 100 μ m) have to be 20 brought to thermal equilibrium. The PCR chamber **731** is preferably situated in a housing with plane-parallel outer surfaces (plastic body **705** with cutout and adhesive film **709**).

In the cartridge receptacle **103**, two thermal coupling 25 plates **733** and **733'**, for example composed of aluminum, are arranged above and below the PCR chamber **731** of the cartridge **101**. The two thermal coupling plates **733** and **733'** are pressed against the adhesive film **709** and against the plastic body **705**, respectively, and serve for transmitting 30 thermal energy of two Peltier units **735** and **735'** into the PCR chamber **731** when a PCR is carried out. The Peltier units **735** and **735'** are in thermal contact with two heat sinks **737** and **737'** which serve for dissipating heat during the cooling operations of the PCR cycles. They have a largest 35 possible surface area in order to enable an efficient heat transfer to the ambient air, if appropriate with the aid of fans.

A cutout **739** is shaped in the thermal coupling plate **733**. By virtue of the cutout **739**, the adhesive film **709** can bulge out in the event of possible excess pressure, for example 40 during a temperature increase in a PCR cycle.

Two permanent magnets **741** and **741'** are arranged centrally above and respectively below the heat sinks **737** and **737'**. The magnetic field generated by the permanent magnets **737** and **737'** serves for holding back magnetic beads 45 when the latter are flushed through the channel **707''** into the PCR chamber **731**. The DNA bound to the magnetic beads is thus held back for later replication by the PCR in the PCR chamber **731**.

The field gradient generated by the two permanent mag- ⁵⁰ nets **741** and **741'** is too small, however, to efficiently hold back the magnetic beads. In order nevertheless to obtain a high field gradient in the PCR chamber **731** along the throughflow direction of the liquid containing the magnetic beads, the thermally conductive but magnetically neutral ⁵⁵ thermal coupling plates **733** and **733'** are equipped with small-volume bodies **743** and **743'** (approximately 5 mm³) composed of a material having a high relative permeability $\mu(r)$, in conjunction with still good thermal conductivity, preferably permalloy (Ni—Fe) or Mumetal (Ni/Fe/Cu/Mo). 60

The small-volume bodies **743** and **743'** respectively form a magnetic core directly at the surfaces of the PCR chamber **731**. The PCR chamber **731** is then situated directly between the magnetic cores. The magnetic cores are magnetized by the magnetic field of the permanent magnets **741** and **741'**. 65 The magnetic field lines are concentrated by way of the magnetic cores, whereby a high field gradient is generated.

In the field gradient arranged parallel to the outer walls of the PCR chamber **731** and flow direction of the solution containing magnetic beads, the magnetic beads are drawn in, detained and therefore concentrated.

The PCR chamber **731** can be closed up by way of two plungers **745** and **745**'. The plungers **745** and **745**' can be pressed against the adhesive film **709** by way of springs and cams that are not illustrated here, such that the PCR chamber **731** is closed off.

The heat sinks **737** and **737'** are connected to screws **761** via springs **759**. The screws **761** are screwed into the cartridge receptacle **103**. The thermal coupling plates **733** and **733'** connected to the heat sinks **737** and **737'**, respectively, are pressed against the adhesive film **709** and the plastic body **705**, respectively, by the springs **759**, such that good thermal contact arises. At the same time, thermal contact between the thermal coupling plates **733** and **733'** and the cartridge receptacle **103** is avoided to the greatest possible extent.

FIG. 11 illustrates a section through a part of the cartridge receptacle 103 with inserted cartridge 101. A further channel 707^{III} is milled into the plastic body 705, said further channel being covered with the adhesive film 709. A chip module 747, by means of which the DNA contained in the sample can be detected, is arranged in the plastic body 705. By way of example, a redox cycling process for detecting the DNA can take place on the chip module 747.

In order to support the hybridization between the DNA and corresponding capture molecules that takes place at the surface of the chip module **747**, the chip module **747** is thermally coupled to a Peltier element **753** via a thermal coupling plate **751**. The Peltier element **753** is thermally coupled to a cooling unit **755**. The temperature in the vicinity of the chip module **747** can be monitored by way of a temperature sensor **757** arranged in the thermal coupling plate **751**. SNP analyses, for example, can thus be carried out by means of determining melting curves.

Outside the plane of the drawing, the unit formed from the thermal coupling plate **751**, the Peltier element **753** and the cooling unit **755** is connected to the lower part **719** of the cartridge receptacle. In this case, the unit is mounted in resilient fashion, such that it is pressed against the cartridge **101** and a good thermal contact between the thermal coupling plate **751** and the chip module **747** is established.

Moreover, outside the plane of the drawing, sprung contact pins are provided within the lower part **719** of the cartridge receptacle **103**, and tap off electrical signals of the chip module **747** for evaluation.

A practically suitable determination of DNA, proteins and also haptens is now possible with the described system including smart card for sample uptake and evaluation device with control device for sample processing and evaluation when the cartridge has been introduced into the evaluation device.

Example embodiments being thus described, it will be obvious that the same may be varied in many ways. Such variations are not to be regarded as a departure from the spirit and scope of the present invention, and all such modifications as would be obvious to one skilled in the art are intended to be included within the scope of the following claims.

What is claimed is:

1. A method for operating a control device in a biochip analysis device comprising:

a cartridge in the form of a plastic card which contains a system of microfluidic channels that form a reservoir that contains a reaction substance, a sample chamber

and flow channels, a fluid input and a deformable means for covering the system of microfluidic channels.

- a control device having a cartridge receptacle for receiving, locking, unlocking and releasing the cartridge, the 5 control device comprising:
 - a power supply connected to the cartridge receptacle. at least one first sensor adapted for identifying whether
 - or not a cartridge located in the cartridge receptacle is a correct cartridge or correctly located therein,
 - a microcontroller adapted for control and data acquisition, and having means for transmitting data to an evaluation unit or computer, and
 - a pump controlled by the microcontroller and adapted to direct a flow of liquid through a cartridge located in the cartridge receptacle,
 - wherein the cartridge receptacle is comprised of upper and lower parts that are movable apart for insertion of a cartridge and together for locking an inserted 20 cartridge in the cartridge receptacle, the method comprising the steps of:

introducing a sample into said cartridge;

- opening the cartridge receptacle;
- using a means for moving to move a part of the cartridge 25 receptacle upon insertion of the cartridge into the cartridge receptacle, such that the cartridge is correctly received in the cartridge receptacle;
- transferring the sample to a PCR chamber;
- dissolving dry PCR reagents;
- closing off valves of the PCR chamber;
- starting thermocycling;
- carrying out a set number of thermocycles at defined temperatures;
- of the PCR;
- transferring the PCR product from the PCR chamber to a detection chamber;
- carrying out an DNA analysis of constituents to be detected in the sample in an integrated process; the 40 carrying out the DNA analysis comprising actively changing the temperature in the detection chamber, such that hybridization of the PCR product with catcher molecules can take place;

pumping dissolved enzyme into the detection chamber; pumping dissolved substrate into the detection chamber; starting electrical detection; and

evaluating measurement signals resulting from the analysis or transmitting data to an external device or computer.

2. The method as claimed in claim 1, wherein an SNP analysis is carried out by recording melting curves.

3. The method as claimed in claim 1, wherein the control device comprises said means for moving a part of the cartridge receptacle and further comprising a means for 55 analysis is carried out by recording melting curves. locking a cartridge inserted into the cartridge receptacle.

4. Method as claimed in claim 1, wherein the control device comprises fill level sensors adapted for monitoring of the flow of fluid in the cartridge.

5. Method as claimed in claim 1, wherein a thermostatic 60 regulation device is provided that has thermal coupling plates at positions adapted for transferring heat to a cartridge in the cartridge receptacle, one of the thermal coupling plates on a side facing the film of an inserted cartridge.

6. Method as claimed in claim 1, further comprising 65 bringing two parts of the cartridge receptacle together by said means for moving.

7. Method as claimed in claim 1, further comprising bringing a Peltier element into position by said means for moving.

8. Method as claimed in claim 1, further comprising performing redox cycling in the detection chamber for detection of DNA.

9. Method as claimed in claim 1, wherein the closing of the valves of the PCR chamber is performed by actuators pressing onto the deformable means for covering the system of microfluidic channels of the cartridge.

10. A method for controlling an integrated and automated analysis of DNA or protein in a system comprising a cartridge containing various reagents for the analysis and to receive a sample to be analyzed and comprising a control device for controlling an analysis process, the method comprising:

introducing a sample into the cartridge;

determining, using a capacitive filling level sensor, a liquid level in a channel of the cartridge;

- actively influencing the temperature in a detection chamber of the cartridge by a Peltier element of the control device which is thermally coupled to a chip module;
- carrying out an analysis of constituents to be detected in the sample in an integrated process;
- recording measurement data from the analysis process via a computing unit;
- transmitting the measurement data to an evaluation unit via a wireless module; and

evaluating the measurement data; and

representing the results of the analysis on a display,

further comprising using a means for moving for bringing the Peltier element into position for said influencing of the temperature in the detection chamber.

11. Method according to claim 10, further comprising: opening the valves of the PCR chamber after conclusion 35 controlling all components of the control device involved in the analysis process via a microcontroller.

> 12. Method according to claim 11, wherein the microcontroller is connected to the computing unit via a serial interface.

> 13. Method according to claim 10, further comprising: controlling a pump for pumping a fluid through channels of the cartridge.

14. Method according to claim 10, further comprising: controlling at least one valve arranged in a channel of the 45 cartridge.

15. Method according to claim 10, further comprising: controlling valves of a PCR chamber via a servomotor and a signal processing unit.

16. Method according to claim 10, further comprising: 50 monitoring the movement of a fluid in the cartridge.

17. Method according to claim 10, wherein carrying out the analysis includes performing a DNA analysis, and the DNA analysis includes cell disruption and PCR.

18. Method as claimed in claim 17, wherein an SNP

19. Method according to claim 10, further comprising: carrying out a thermocycling process for PCR, wherein dry PCR reagents are dissolved.

20. Method as claimed in claim 10, wherein a protein analysis is carried out.

21. Method as claimed in claim 10, further comprising closing valves of a PCR chamber by actuators pressing onto a deformable means for covering the cartridge.

22. Method as claimed in claim 10, further comprising actively influencing the temperature in the detection chamber by a Peltier element which is thermally coupled to a chip module in order to support a hybridization between the replicated DNA and corresponding capture molecules that takes place at a surface of a chip module of the cartridge.

23. A method for controlling an integrated and automated analysis of DNA or protein in a system comprising a cartridge containing various reagents for the analysis and to 5 receive a sample to be analyzed and comprising a control device for controlling an analysis process, the method comprising:

introducing a sample into the cartridge;

- carrying out an analysis of constituents to be detected in 10 the sample in an integrated process;
- closing off a PCR chamber of the cartridge by closing two valves by actuators pressing onto a film of the cartridge;
- pressing a device for transferring heat to the PCR cham- 15 ber against a plastic body of the cartridge when a PCR is carried out, wherein said device comprises two Peltier units which are arranged on opposite sides of the PCR chamber;
- carrying out a thermocycling process for PCR, wherein 20 dry PCR reagents which are stored in the cartridge are dissolved;
- carrying out a predetermined number of thermocycles at defined temperatures;
- recording measurement data from the analysis process via 25 a computing unit;
- transmitting the measurement data to an evaluation unit via a wireless module; and evaluating the measurement data,
- further comprising using a means for moving to bring the 30 device for transferring heat into position against the plastic body.

24. Method as claimed in claim 23, further comprising inserting the cartridge into a cartridge receptacle and bringing two parts of the cartridge receptacle together by said 35 means for moving.

25. Method as claimed in claim **23**, further comprising actively influencing the temperature in a detection chamber of the cartridge by a Peltier element which is thermally coupled to a chip module in order to support a hybridization 40 between replicated DNA and corresponding capture molecules that takes place at a surface of a chip module of the cartridge.

26. Method as claimed in claim **25**, further comprising performing redox cycling in the detection chamber for 45 detection of DNA.

27. Method as claimed in claim **23**, further comprising electrically detecting the PCR product by means of a sensor array.

28. A method for controlling an integrated and automated 50 analysis of DNA in a system comprising a cartridge containing various reagents for analysis and receiving a sample to be analyzed and comprising a control device for controlling an analysis process and having Peltier elements, the method comprising:

introducing a sample to be analyzed into the cartridge; opening a cartridge receptacle of the control device;

- using a motor means for moving a part of a cartridge receptacle upon insertion of the cartridge and/or bringing two parts of the cartridge receptacle together when the cartridge has been completely introduced;
- upon insertion of the cartridge, bringing Peltier elements of the control device into position by another motor means;
- transferring a sample to be analyzed to a PCR chamber of the cartridge;
- closing off the PCR chamber using actuators by pressing two valves onto a film of the cartridge;
- performing PCR by carrying out a predetermined number of thermocycles;
- opening the PCR chamber after conclusion of the PCR; transferring the PCR product from the PCR chamber to a detection chamber of the cartridge;
- actively influencing the temperature in the detection chamber using a Peltier element which is thermally coupled to a chip module in order to support hybridization between the PCR product and corresponding capture molecules that takes place at a surface of the chip module of the cartridge;
- performing redox cycling in the detection chamber for detection of DNA;
- electrically detecting the PCR product by means of a sensor array;
- recording measurement data from the analysis process with a computing unit;

evaluating the measurement data; and

representing the results of the analysis on a display.

29. Method as claimed in claim **28**, further comprising transmitting the measurement data to an evaluation unit via a wireless module.

30. Method as claimed in claim **28**, further comprising interrupting the movement of the cartridge by a limit switch when the cartridge has been completely introduced.

31. Method as claimed in claim **28**, further comprising marking DNA linked to catcher molecules by a label enzyme.

32. Method as claimed in claim **28**, further comprising pressing Peltier units, which are arranged on opposite sides of the PCR chamber for transferring heat to the PCR chamber, against a plastic body of the cartridge when PCR is carried out.

33. Method as claimed in claim **28**, wherein a Peltier element for actively influencing the temperature in the detection chamber is pressed against the cartridge.

34. Method as claimed in claim **28**, wherein the sensor array comprises a CMOS sensor chip.

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