

# Direct analysis of dried blood spots coupled with mass spectrometry: concepts and biomedical applications

Julien Déglon · Aurélien Thomas · Patrice Mangin ·  
Christian Staub

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**Abstract** Because of the emergence of dried blood spots (DBS) as an attractive alternative to conventional venous plasma sampling in many pharmaceutical companies and clinical laboratories, different analytical approaches have been developed to enable automated handling of DBS samples without any pretreatment. Associated with selective and sensitive MS–MS detection, these procedures give good results in the rapid identification and quantification of drugs (generally less than 3 min total run time), which is desirable because of the high throughput requirements of analytical laboratories. The objective of this review is to describe the analytical concepts of current direct DBS techniques and to present their advantages and disadvantages, with particular focus on automation capacity and commercial availability. Finally, an overview of the different biomedical applications in which these concepts could be of major interest will be presented.

**Keywords** Dried blood spots · Direct LC–MS–MS analysis · Ambient MS · Automated on-line DBS

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J. Déglon · A. Thomas · P. Mangin · C. Staub (✉)  
Unit of Toxicology, CURML,  
1 rue Michel-Servet,  
1211 Geneva 4, Switzerland  
e-mail: Christian.staub@hcuge.ch

J. Déglon · A. Thomas · C. Staub  
Swiss Centre for Applied Human Toxicology,  
University of Geneva,  
1211 Geneva, Switzerland

## Introduction

Although biological analysis is usually performed using plasma obtained by venipuncture, the use of dried blood spots (DBS) has grown in popularity in the clinical and pharmaceutical communities over the past decade as an alternative sampling procedure. The numerous advantages of the DBS sampling process can be summarized as follows:

- **Ease** – DBS is less invasive than conventional whole blood, plasma, or serum sample collection [1–3] because the blood can be collected after a small finger prick (or heel prick for pediatric applications). Because of the ease of collection, DBS can be obtained in a non-hospital environment by minimally trained technicians or even at home by the patients themselves [4–6]. Moreover, the DBS process does not require use of an anticoagulant, or plasma separation, which limits the number of manipulations.
- **Small sampling volume** – less than 20 µL is usually spotted on to filter paper, whereas a minimum volume of 0.5 mL is required with conventional venous sampling.
- **Cost-effective process** – the cost of shipping/storage of filter paper cards is substantially reduced, because the DBS samples can be stored at room temperature without the use of dry ice [7].
- **Analyte stability** – it has been reported in numerous publications that DBS that have been stored over many months (even years) at ambient temperature are just as stable as plasma samples that were stored at –20 °C [8, 9].
- **Improved safety** – during drying, most of the pathogenic agents are deactivated on the filter paper, which reduces the risk of infection to a minimum. Similarly, transfer of blood material on to a filter paper card is considered to have a low infection risk [4, 10, 11].

These advantages make the DBS procedure a patient-friendly tool for blood collection, especially for problematic and vulnerable patient populations [12–14]. The ease of the process can also help in the recruitment of subjects (human and animal) for preclinical and clinical studies [15–17] and can significantly reduce the cost of these studies [18]. Interesting reviews covering these different aspects were recently published to promote the use of DBS in bioanalysis [19, 20].

The first biomedical application of DBS on filter paper dates back to 1963 when Professor Robert Guthrie introduced this alternative sampling method for detection of phenylketonuria in the newborn population [21]. Detection of L-phenylalanine was based on a microbiological test that was sufficiently sensitive but with low analytical throughput. In the early 1990s, the development of PCR and immunoassays, including ELISA, RIA, or FIA, enabled the detection of DNA, viral RNA, antibodies, and hormones from DBS with an acceptable waiting time that was suitable for high-throughput analyses [22–27]. These advances extended the scope of applications of the DBS to the detection of different biomarkers in epidemiological studies [11, 28].

At the same time, the detection of small pharmaceutical compounds was investigated by coupling liquid chromatography (LC), as separation technique, with UV or fluorescence detection. Although LC was, in general, more selective than immunoassays by separating the compounds of interest from the matrix before detection, the initial methods based on DBS–LC–UV were limited to the quantification of drugs at high concentrations [29–31], because of the poor sensitivity.

In fact, one limitation of DBS analysis is that the small volume of biomatrices spotted implies the need for highly sensitive techniques for detection and quantification of molecules present at concentrations below  $1 \text{ ng mL}^{-1}$  [19]. A new generation of mass spectrometers (MS), particularly triple-quadrupole spectrometers (MS–MS) operating in MRM mode, have greatly improved sensitivity, which enables analysis of pharmaceuticals in the low therapeutic range [32]. Therefore, many studies have recently been published showing interest in coupling DBS samples with LC–MS–MS for measurement of therapeutic agents or biomarkers in clinical and research settings [33–37].

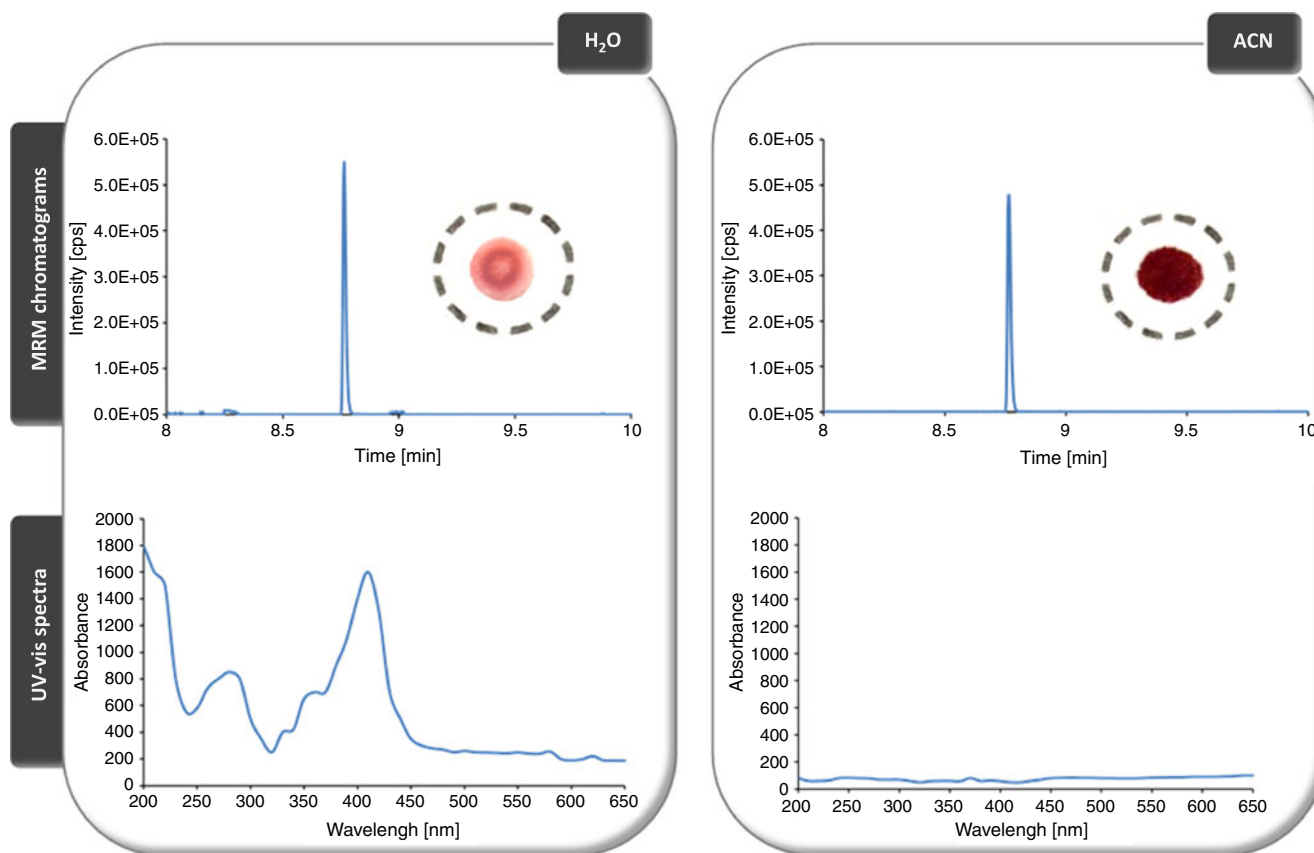
Before detection, the analytes must be extracted from the DBS to be more compatible with the analytical technique (for example, immunoassays or LC–MS–MS). Thus, an extraction is usually performed in off-line mode using a mixture of aqueous and organic solvents. This pretreatment step requires time and manipulation by technicians and can become problematic if the number of samples is large. Interestingly, the implementation of DBS support in pharmaceutical companies in the last five years has led to

the development of novel techniques for handling this new support before the analysis. This review will, therefore, highlight the recent developments that enable direct analysis of DBS samples either by direct insertion into conventional LC–MS–MS systems or by direct ambient extraction/ionization with subsequent MS–MS detection. Finally, an overview of the different biomedical applications will be presented.

## Analytical concepts

### On-line DBS

Integration of DBS with LC–MS is generally performed by solvent extraction, which ensures transfer of the analytes from the filter paper into an injectable solution that is compatible with LC, as discussed above. Emphasis on DBS sampling, especially in pharmaceutical analysis, has led to the development of novel techniques that enable high-throughput analysis. In a recent report, Déglon et al. described a novel concept that enabled on-line desorption of filter paper into an LC–MS system without any sample pretreatment [38]. For this purpose, a stainless steel desorption cell compatible with LC capillaries was designed to integrate the filter paper containing the dried spot. The behavior of the DBS during dynamic desorption (ensured by application of a constant flow rate) was also investigated to determine the kinetics and the efficiency of analyte desorption and their dependence on experimental conditions such as solvent and the flow rate. Experiments have shown that the type of solvent can selectively affect desorption output. As shown in Fig. 1, use of an aqueous solution led to complete desorption of analytes and matrix components, for example hemoglobin and proteins, which were monitored using MS–MS and UV–visible spectroscopy, respectively. Interestingly, use of organic solvents such as ACN or MeOH led to contrasting results—only the analyte of interest was monitored and the matrix components were not detected. This was in agreement with the color of the filter paper, which was maintained after organic desorption. In another study, Thomas et al. showed that selective desorption was applicable to a wide range of polarity compounds, for example hydrophobic and hydrophilic drugs and their phase I and II metabolites [15]. These results demonstrated that filter paper can not only be regarded as a biofluid support. Indeed, the intrinsic properties of cellulose have to be considered when developing sample preparation or investigating novel approaches, for example the development of selective chemical or affinity interactions on the filter paper. In this case, percolation of organic solvents through the filter paper induced protein precipitation “on support,” which enabled selective desorption of small molecules, for example drugs



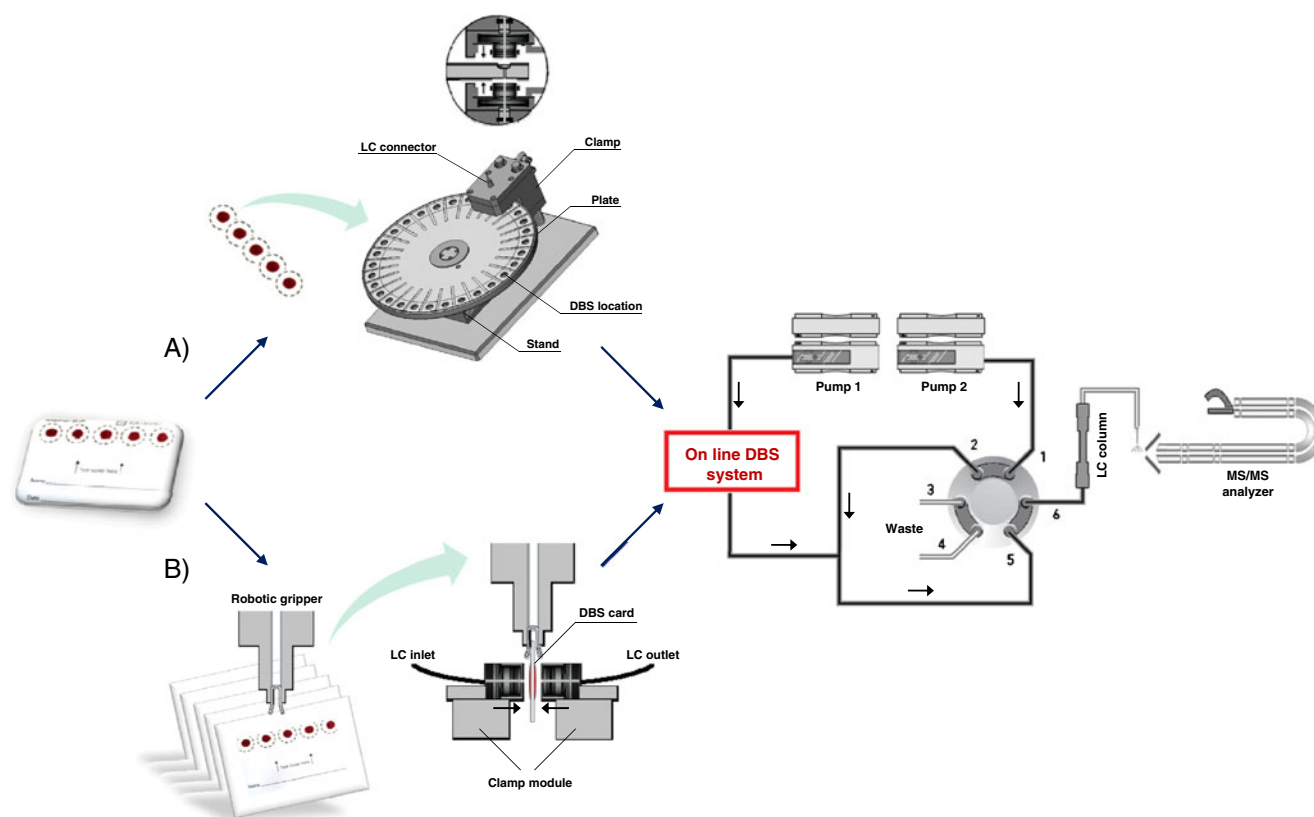
**Fig. 1** MRM chromatograms (*top*) and associated UV–visible spectra (*bottom*) obtained after on-line extraction with H<sub>2</sub>O (left) and ACN (right) of DBS samples spiked with saquinavir at 250 ng mL<sup>-1</sup>. Pictures of DBS samples after extraction are incorporated in the MRM chromatograms

and pharmaceutical compounds. Because of the well-known ion suppression encountered with atmospheric pressure ionization sources such as ESI, the ability to eliminate matrix effects without any sample preparation can be particularly beneficial when LC–MS analysis is performed. Post-column infusions have confirmed that this experimental design is promising in reducing or avoiding matrix effects, compared with conventional analysis of plasma. As illustrated in Fig. 1, use of an aqueous desorption mobile phase (MP) can lead to rapid clogging of analytical systems, the risk of matrix effects, and the need to change chromatographic columns frequently.

The kinetics of compound desorption have been investigated, and they were dependent on the flow rate and cell geometry. In the first test of cell geometry (in which a 10 mm i.d. filter paper was used), a flow rate of 0.25 mL min<sup>-1</sup> was shown to give complete desorption after 5 min. Reduction of dead volume is particularly important to reduce desorption time to a few seconds while maintaining a low flow rate (0.1 mL min<sup>-1</sup>). However, this reduction implies a concomitant decrease in the size of the DBS used, and therefore in sensitivity, because, proportionally, a lower volume of blood will be desorbed.

The fact that desorption was performed with an organic solvent required the polarity of the MP to be adjusted using an additional pump (delivering aqueous MP) after desorption to ensure good trapping of the analyte on the reversed-phase column and to avoid compound dispersion. Therefore, focusing of the compounds can be performed on a trapping column in a column-switching setup or directly on the analytical column (because of very low desorption of the contaminants).

This concept was recently automated by Déglon et al. [39]. As illustrated in Fig. 2A, the developed prototype was based on an automated rotating plate machined to host multiple wells in which filter paper can be placed manually. Next, an automatic clamp that contains two pistons was programmed to sequentially ensure locking of the well positions (followed by the desorption step) and was synchronized with an LC–MS–MS system. This system setup enabled many DBS samples to be analyzed. Furthermore, Thomas et al. demonstrated that reduction of the flow rate to a few  $\mu\text{L min}^{-1}$  with the automated system gave a desorption profile that was very close to an infusion profile, which enabled MS acquisition over several minutes [40]. Based on the selectivity of the process discussed above, this



**Fig. 2** Illustration of two automated on-line DBS instruments, the automated on-line DBS prototype developed by Déglon et al. (A) and the SCAP solution commercialized by Prolab (B), and their respective integration into conventional LC–MS–MS systems

approach was applied in a lipidomic approach for direct MS analysis of dried spots without any sample preparation or chromatographic separation (see the section “[Biomarkers identification](#)”).

In parallel, the robot industry has recently developed automatic systems for on-line DBS analyses [41, 42]. Prolab Instruments recently presented a commercially available solution called SCAP DBS based on the principles discussed above. In their configuration, an automatic clamp (with different available i.d. sizes) enables direct desorption of DBS integrated to a column-switching system with the presence of an auxiliary pump for dilution of the desorption MP [43]. The setup works directly on the DBS card without the previous punch-out step because desorption of the blood spot is ensured by the on-spot locking of the clamp, as shown in Fig. 2B. Furthermore, this system is combined with the versatile CombiPal autosampler for successive handling of multiple DBS cards. In this configuration, a desorption of 0.5 min (for a total run time of less than 4 min) was shown to be a good compromise between sensitivity and extraction efficiency.

Compared with off-line extraction, which dilutes the analyte, a remarkable advantage of on-line DBS procedures is the efficient desorption of the compounds of interest

directly into the analytical systems; it is therefore a non-diluting procedure [44]. Recent applications based on automated on-line DBS analysis coupled to LC–MS–MS systems (in MRM mode) have enabled full validation of this technique using different pharmaceutical probes, for example oseltamivir [43], flurbiprofen and its phase I metabolite [39], and six tyrosine kinase inhibitors (data not given). Good accuracy and precision (<15%) have been obtained for these compounds, and the limit of quantification is as low as  $1 \text{ ng mL}^{-1}$ . For these automatic systems carryover is low; this is explained by the cleanup position of the clamp after DBS extraction. The different aspects of the techniques that were discussed in this section are summarized in Table 1.

### Surface sampling probe

Compared with the on-line DBS approach in which the analytes are desorbed from the filter paper by percolation of a solvent through the blood spots, a second approach, called surface sampling probe (SSP), extracts the compounds of interest from a surface by connecting this surface with a confined liquid stream dispensed by an appropriate probe [45]. The stream is both brought to the surface then carried on to the conventional LC–MS device or directly infused

**Table 1** Evaluation of the different techniques that enable direct DBS analysis coupled with MS–MS detection

Criteria	Direct DBS analysis techniques				
	Online DBS (SCAP)	SSSP (TLC–MS)	LMJ-SSP (LESA)	Paper spray	DESI (Omni Spray)
LC separation-compatible	Yes	Yes	No	No	No
Sensitivity	****	****	**	**	**
Quantitative performance	****	****	**	*	*
Automation/throughput	***	* (***) <sup>a</sup>	****	*	*
Commercial availability	Yes	Yes	Yes	No	Yes
Cost	**	*	***	n.d.	*

<sup>a</sup> Available soon

Criteria evaluation is based on four levels corresponding to low (\*), medium (\*\*), high (\*\*\*), and very high (\*\*\*\*). For cost evaluation, one \* corresponds to a price going from 10,000 to 50,000 \$.

into the MS ionization source. Two different types of SSP have been published—the liquid microjunction-surface sampling probe (LMJ-SSP), which can be applied to extraction of analytes from the surfaces of hard and nonporous materials like glass or metal, and the sealing surface sampling probe (SSSP), by use of which porous surfaces, including most filter papers, can also be analyzed.

### SSSP

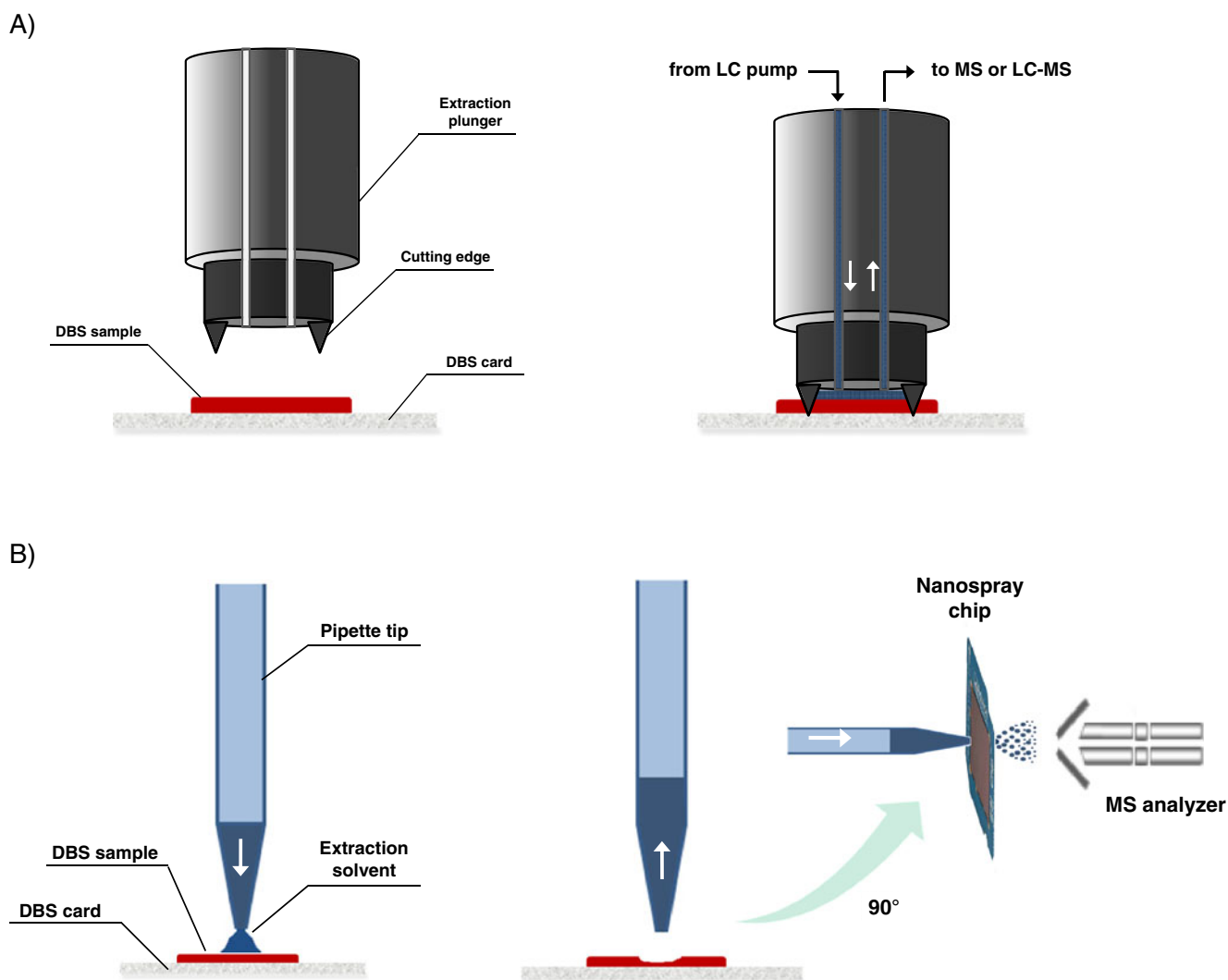
Direct liquid extraction based on SSSP is part of an emerging area in MS that combines ambient extraction of analytes contained in or on surfaces with ESI or APCI ionization. Initially, SSSP was developed to couple cost-effective thin layer chromatographic (TLC) separation with selective and sensitive MS detection. Luftmann reported an original device that enabled on-line transfer of TLC-separated compounds into a solvent stream toward MS [46]. Based on this proof-of-concept, Camag commercialized a TLC–MS interface that had the same characteristics as the Luftmann plunger [47]. However, unlike the latest development of the Luftmann plunger in terms of automation [48], Camag did not include automatic positioning of the plunger above the sample foil in its first version.

More recently, Van Berkel et al. extended the original use of the Camag TLC–MS interface for DBS analysis by replacing the TLC plates with filter paper sheets containing DBS samples [45]. Briefly, the Camag interface used for this approach comprised a stainless steel plunger (also called an extraction head) with a solvent inlet and an outlet capillary and a ring-shaped cutting edge at its end face (Fig. 3A). Before analysis, the DBS samples were manually positioned under the extraction head assisted by a laser pointer, and the plunger was activated and pressed on the paper sheet by compressed air. The cutting edge severed the filter paper and formed a tight seal with the paper. The extraction solvent, which was delivered by a conventional

LC pump, entered the plunger through the inlet capillary, dissolved the analytes contained in the DBS, and left through the outlet capillary before ESI–MS detection. To avoid any clogging of the instrument, the outlet capillary was protected by a filter frit. A six-port valve enabled isolation of the solvent flow from the MS interface when the plunger was in the retracted position. Thus, the capacity of the Camag TLC–MS equipment to directly extract drugs from DBS previously spotted on filter paper was demonstrated for the first time. For these tests, sitamaquine and acetaminophen were used as probes to evaluate the reproducibility and the quantitative aspects of the system. Experiments were conducted using MeOH as extraction solvent at a flow rate of 0.2 mL min<sup>−1</sup> through a 4 mm i.d. plunger for a duration of 60 s, and the extraction was followed by direct MS detection in MRM mode. To eliminate carryover, the system was washed for 60 s between each DBS by sampling a blank zone on the filter paper. The spot-to-spot analysis time was approximately 3 min, which includes the DBS extraction and cleanup steps. The reproducibility of the extraction was excellent, carryover was low when using washout, and good quantification was achieved over the concentration ranges 10 to 10,000 ng mL<sup>−1</sup> and 50 to 5000 ng mL<sup>−1</sup> for sitamaquine and acetaminophen, respectively. Unfortunately, the effect of the matrix was not investigated in this study.

In parallel studies, Abu-Rabie et al. tested the Camag TLC–MS interface in combination with LC separation before MS detection [49]. On the basis of the same LC–ESI–MS–MS method, they compared traditional off-line DBS extraction and both modes of the Camag TLC–MS (with or without the LC column). For these experiments, different solvents, flow rates, and extraction times were tested in the analysis of DBS for various pharmaceuticals. Extraction solvents giving good sensitivity were usually a mixture of aqueous buffer and MeOH or ACN. Because of the lack of a second LC pump that could deliver H<sub>2</sub>O to





**Fig. 3** Schematic views of the extraction head of the Camag TLC-MS interface (**A**) and the TriVersa NanoMate of Advion operating in LESA mode (**B**) for the direct extraction of DBS samples with subsequent MS detection

focus the analytes ahead the analytical column, a higher flow rate was applied (between 0.5 and 1 mL min<sup>-1</sup>) during a very short period to obtain sufficient extraction efficiency. Under these conditions, the chromatographic peak shape was acceptable up to an extraction time of 5 s, after which it became increasingly broad. Use of short (i.e. 50 mm) LC columns with medium-sized (3 µm) particles enabled the application of high flow rates without generating excessive backpressure in the plunger. Despite the use of a very short extraction period, improved sensitivity was achieved compared with the off-line extraction mode, with which only part of the extraction solvent could be injected, which implies an important dilution factor. Bioanalytical validation of the TLC-MS interface in LC-MS-MS mode was performed on sitamaquine and acetaminophen. According to international validation criteria, acceptable accuracy, precision, and linearity were achieved over the ranges 5 to

1000 ng mL<sup>-1</sup> and 50 to 50,000 ng mL<sup>-1</sup> for sitamaquine and acetaminophen, respectively. However, a substantial amount of carryover was observed; this required manual cleaning of the system.

Subsequently, Heinig et al. presented a TLC-MS interface that was integrated into a column-switching LC-MS-MS system that enabled quantification of drugs in DBS at concentrations as low as 50 pg mL<sup>-1</sup> [44]. The increase in sensitivity compared with the work discussed above was because of the combination of a longer extraction step (typically 45 s) with focusing of analytes ahead of the trapping column ensured by a tee connector and a second LC pump delivering high H<sub>2</sub>O flow rate (see the section “On-line DBS”).

The SSSP technique, by means of the commercial TLC-MS interface, has proved to be a promising method with sufficiently reproducibility and sensitivity with a suitable

delay for high-throughput DBS analysis (Table 1). Intrinsically, the total run time using LC separation was similar to the run times obtained by use of SCAP technology [43]. However, the TLC–MS interface suffers from an important lack of automation, because three manual operations (DBS positioning below the extraction head, activation of the plunger, and the six-pore valve switch) were required before the start of an analysis. To handle larger sample numbers by feeding the DBS card automatically into the extraction device, Camag is currently developing an automated system for routine application which should be commercially available during 2011 [50].

### LMJ-SSP

In the LMJ-SSP approach, the analytes are extracted from the surface by connecting the surface to a wall-less liquid microjunction dispensed by an appropriate probe. Unlike the SSSP technique, in which the plunger forms a sealed contact with the sampling surface, the probe used in the LMJ-SSP is positioned an appropriate distance from the surface (typically 0.2 mm), and a liquid microjunction is then formed by enabling the liquid from the end of the probe to extend from the probe to the surface [51]. Because of the dynamic solvent flow and the lack of sealing, the current LMJ-SSP probe is unsuitable for extraction of porous surfaces. In the analysis of DBS, the liquid microjunction will not remain in contact with the filter paper because of its high absorption capacity. However, an interesting alternative which overcomes this limitation is use of the Advion TriVersa NanoMate system operating in LESA mode in which a static microjunction can be performed. As depicted in Fig. 3B, the TriVersa NanoMate is constituted of two distinct parts: a robotic arm with adequate conductive pipette tips ensuring that samples are loaded, and a chip that contains microfabricated nozzles to generate a nanoelectrospray of the liquid samples. Before the surface sampling process, the robotic arm picks up a conductive pipette tip and moves above the well that contains the extraction solvent. A few microliters (usually 1 to 5  $\mu\text{L}$ ) are loaded in the pipette tip and the arm automatically goes to the desired extraction zone (DBS). The arm is then lowered to an appropriate distance from the target spot, and part of the extraction solvent is dispensed on to the sample surface to form a static liquid microjunction between the end of the pipette tip and the surface. Immediately thereafter, the solution is aspirated back into the tip and is finally sprayed through a nanospray nozzle for subsequent MS detection.

Kertesz and Van Berkel published an interesting paper on the feasibility of this concept [52]. Based on the NanoMate TriVersa operating in LESA mode, they performed quantitative analyses of pharmaceuticals from three different

surfaces—MALDI plates, thin tissue sections, and DBS. For the DBS, quantitative investigation was performed using sitamaquine in a 4-mm DBS using a mixture of ACN, MeOH, and an aqueous buffer as the extraction solvent. In this experiment, 4  $\mu\text{L}$  was loaded in the pipette tip and 3  $\mu\text{L}$  was dispensed on to the DBS samples before nanoESI–MS–MS detection; this led to a total run time of approximately 2 min per sample.

Quantitative performance was also investigated by the authors. In accordance with international guidelines, LOQ was set to 100  $\text{ng mL}^{-1}$  (<15% precision, <15% accuracy). An important bias (77%) and poor precision (98%) were observed at lower concentrations (10  $\text{ng mL}^{-1}$ ) even though a deuterated internal standard was used. Quantitatively, on-line DBS or SSSP could reach an LOQ that was 100 times lower [39, 44]. This can be explained by the use of dynamic extraction flow applied to the DBS compared with the static transfer that occurred in the LMJ delivered by the LESA pipette tip. Moreover, the lack of an LC setup to ensure focusing of the desorbed analytes contributes to limiting the sensitivity.

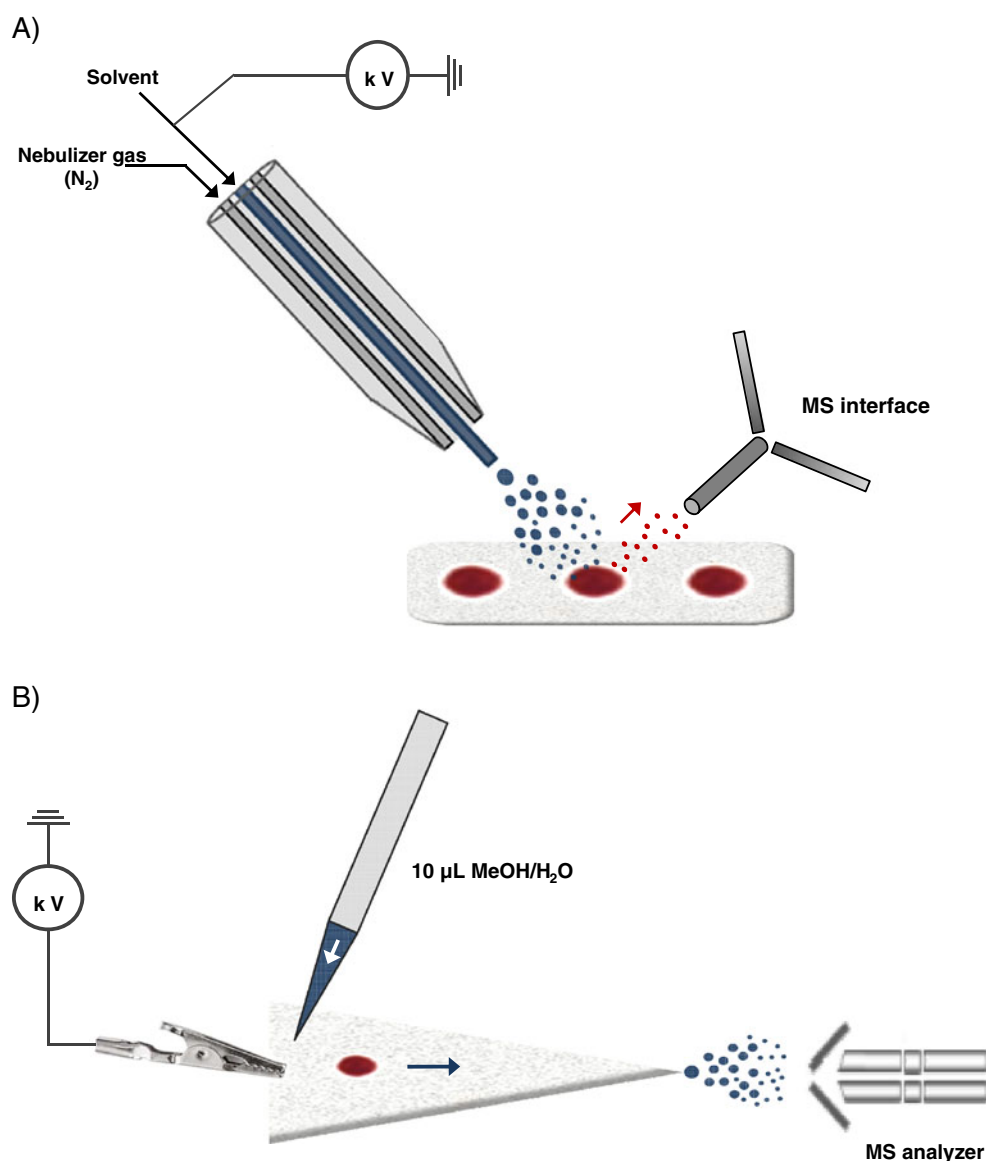
However, LMJ-SSP with the TriVersa NanoMate has two principal advantages compared with the Camag TLC–MS interface. First, it is a carryover-free approach because the pipette tip and nanospray nozzle are changed between each analysis, which means that no cleanup step is required. Second, the procedure is automated, which enables high-throughput analysis.

Although the TriVersa NanoMate combines the advantages of a nanoESI source with direct extraction of a sampling surface, the cost of this system is 10 times higher than the instrument commercialized by Camag (Table 1).

### Desorption electrospray ionization and paper spray

In this section, we will discuss two ambient ionization MS (AIMS) methods used for direct analysis of the DBS—desorption electrospray ionization (DESI) and paper spray (PS). The AIMS techniques enable ionization of untreated samples in an open environment [53]. No sample pretreatment is required and analysis is very fast with the time scale limited to the time needed to present the sample to the MS [54]. Because of these features, AIMS methods are particularly suitable for high-throughput analysis.

Among recent AIMS techniques, DESI developed by Cooks et al. in 2004 is regarded as the initial technique. Briefly, the main DESI mechanism can be described as “droplet pick-up” [55] (Fig. 4A). In this process, ions are generated from the sampling surface by bombardment with charged micro-droplets through the atmosphere. The impact of the spray causes the formation of a microscopic layer of liquid on the sample surface, which results in dissolution of the analytes. This process is directly followed by desorption



**Fig. 4** Schematic representations of the DESI source (A) and the PS configuration (B) applied to direct MS analysis of the DBS samples

by momentum transfer when additional droplets collide with the liquid layer and force the dissolved analytes into the gas phase as micron-sized droplets [56]. After their evaporation, which is assisted by a standard heated capillary extending from the MS inlet, the charged droplets produce free gas-phase ions when entering the MS interface.

Recently, Wiseman et al. experimented with DESI methodology for direct analysis of pharmaceutical compounds in DBS [57]. For this purpose, sitamaquine, terfenadine, and prazosin were chosen as probes and verapamil was used as the common internal standard. In these preliminary experiments, DBS were cut into strips and secured to a glass slide, and a commercial DESI source

(Prosolia Omni Spray 1D Ion Source) was coupled with an MS–MS instrument. Each DBS was therefore analyzed by scanning across the DBS surface at a rate of  $200 \mu\text{m s}^{-1}$ , which resulted in a peak 40 s wide on the ion chromatogram. The solvent flow rate delivered by the sprayer was also investigated and was set at  $15 \mu\text{L min}^{-1}$  as a good compromise between sensitivity and cross-contamination. Good signal intensity and stability were obtained when solvent extraction was performed with 80:20 ACN–H<sub>2</sub>O containing 0.1% formic acid.

Under these conditions, full validation was performed on sitamaquine in the concentration range from 10 to 10,000 ng mL<sup>-1</sup>. Unfortunately, accuracy and precision of the values for QC samples at 10 and 40 ng mL<sup>-1</sup> were



outside the pre-defined performance criteria (>15%). Thus, the quantitative performance of DESI was similar to that of LMJ-SSP using LESA (see the section “LMJ-SSP”). The authors also investigated the ion-suppression effect. They demonstrated that the concentration of IS added (between 500 and 2500 ng mL<sup>-1</sup>) led to suppression of the sitamaquine response by a factor of ten. This ion suppression explains the quantification problems that arose at low concentration.

An alternative to DESI for direct DBS analysis is the PS approach. This technique enables ions to be generated from a liquid sample loaded on to a piece of filter paper by applying a high potential to the wet paper [54]. Indeed, the wet paper is conductive and the network of cellulose fibers forms microchannels for liquid transport. As shown in Fig. 4B, the high potential applied between the base of the paper (previously cut into a triangular shape) and the MS inlet generates an electric field that induces charge accumulation at the apex of the paper triangle, which faces the MS inlet. Similar to ESI, Coulombic forces fracture the liquid to form charged droplets, which undergo desolvation and generate dry ions. On the basis of this mechanism, Liu et al. recently developed a method that enabled direct analysis of atenolol from a 0.4-μL DBS previously spotted on to filter paper [58]. Briefly, triangular paper (10 mm long and 5 mm wide at its base) containing the DBS sample was held by a copper clip in front of the MS inlet at a distance of 3 mm. Next, 10 μL 1:1 MeOH–H<sub>2</sub>O was applied near the base of the paper triangle to wet the paper and extract the analytes, and a 4.5-kV potential was applied while the mass spectra was recorded in MRM mode. The results showed the LOD was 1 μg mL<sup>-1</sup> for atenolol; no other quantitative aspects were investigated. As previously described, use of an organic solvent (MeOH in this instance) enabled the blood cells to remain on the filter paper, whereas dissolved molecules were carried to the paper tip and released in the spray.

Wang et al. showed the quantitative potential of this technique by quantifying imatinib in a 0.4-μL DBS in the concentration range 62.5 to 4000 ng mL<sup>-1</sup>, with labeled compound used as an internal standard [59]. Nevertheless, full validation of this process must still be completed to characterize the technique.

The AIMS methods, for example DESI and PS, are an interesting alternative for direct DBS analysis, because the procedure is relatively user-friendly and pretreatment-free. However, the very small blood volume spotted (0.4 μL for PS) and the lack of LC columns results in relatively low sensitivity compared with other methods (Table 1). Moreover, substantial ion suppression in the DESI approach limits quantitative performance.

## Applications

Although plasma sampling is still the gold standard in pharmaceutical and clinical laboratories, DBS is becoming increasingly accepted in biomedical applications, particularly in pediatrics in the USA, where 95% of newborns are screened for inborn metabolic disorders [60, 61]. As demonstrated by recent publications, this alternative sampling procedure has been successfully introduced for other applications, including clinical investigations, TDM programs, and epidemiological or surveillance studies [62]. Although these methods are mainly based on off-line extraction of filter paper, promotion

**Table 2** Representative MRM transitions of pharmaceuticals and metabolites

Analyte	MRM transition <sup>a</sup> ( <i>m/z</i> )	Estimated LOQ <sup>b</sup> [ng mL <sup>-1</sup> ]
Alprazolam	309→281 (35)	1.0
Amiodarone	646→58 (90)	0.5
Atracurium	358→206 (27)	0.5
Benzoylcegonine	290→168 (29)	1.0
Bupropion	240→131 (30)	0.5
Citalopram	325→109 (20)	1.0
Clopidogrel	322→212 (23)	0.5
Cocaine	304→182 (29)	1.0
Codeine	300→165 (50)	5.0
Dextromethorphan	272→128 (85)	1.0
Diazepam	285→193 (35)	10
Fentanyl	337→188 (35)	0.2
Flunitrazepam	314→268 (35)	5.0
Imatinib	494→394 (36)	1.0
Methadone	310→265 (20)	1.0
Midazolam	326→291 (37)	0.5
Mirtazapine	266→195 (35)	0.5
OH-midazolam	342→324 (31)	2.0
Omeprazol	346→198 (19)	0.5
Phenytoin	253→182 (20)	50
Prasugrel	374→206 (23)	0.2
Quetiapine	384→253 (35)	1.0
Risperidone	411→191 (35)	5.0
Saquinavir	671→570 (45)	2.0
Tacrolimus	821→768 (30)	5.0
Tramadol	264→58 (20)	0.5
Venlafaxine	278→260 (20)	5.0
Zolpidem	308→235 (35)	1.0

<sup>a</sup> The collision energy (eV) is indicated in parentheses

<sup>b</sup> Limits of quantification (*S/N*>10) estimated for 5 μL DBS samples analyzed on a 5500 QTrap mass spectrometer in ESI mode. Extraction was performed by use of automated on-line DBS followed by RP-LC separation on a Kinetex C<sub>18</sub> column (2.1 mm×50 mm, 2.7 μm i.d.)

of the novel techniques discussed in this review will ensure new opportunities in these fields. The next sections of this review will present a non-exhaustive list of applications for which direct and automated DBS analysis are suitable.

### General unknown and target screenings

In biomedical applications in which rapid and comprehensive analytical tools are required, for example in general unknown screening (GUS) and target screening (TS), direct DBS analysis may be an attractive solution because it combines a friendly sampling process with sensitive and selective MS–MS detection.

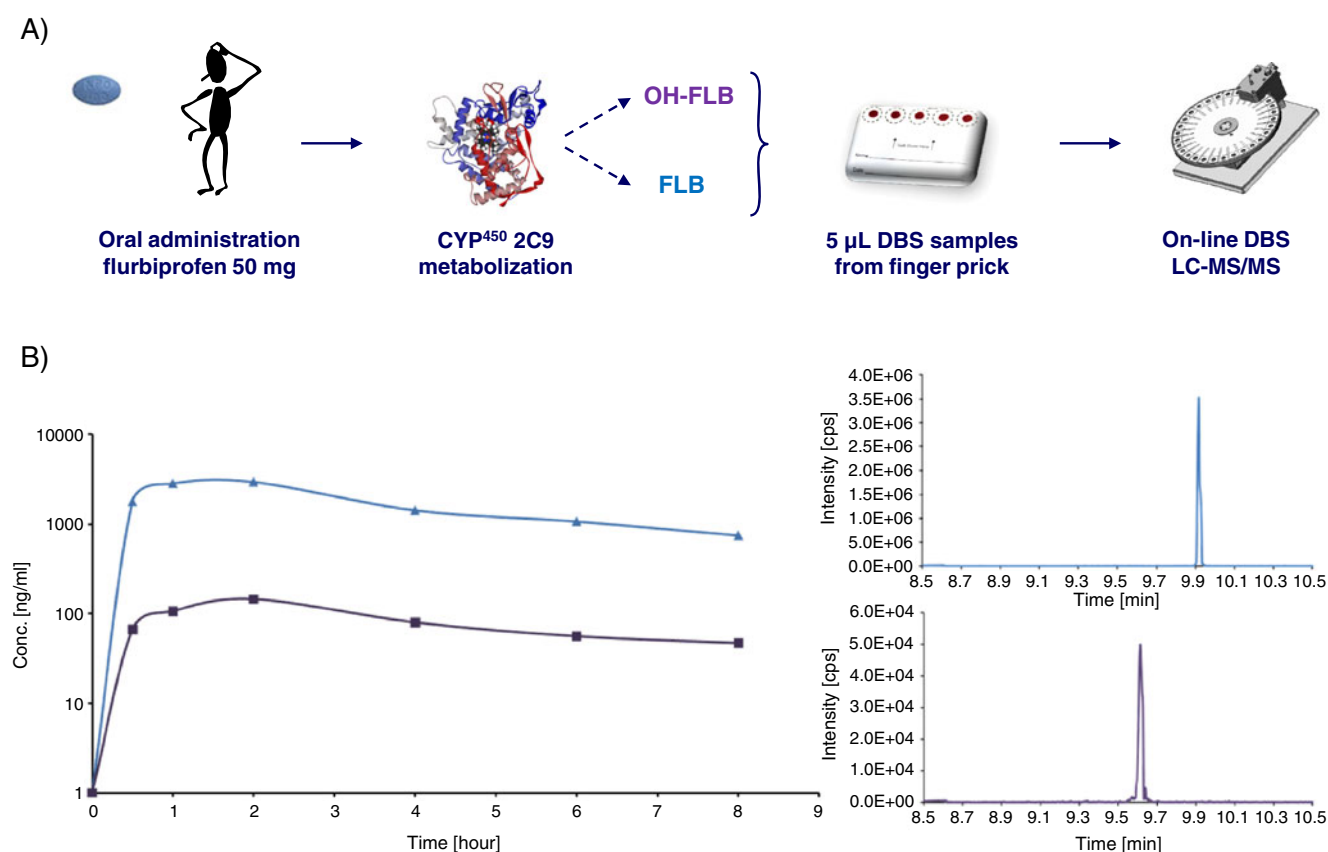
LC–MS–MS has, moreover, recently been applied to GUS and TS strategies [63]; the capacity of direct DBS analysis enabled simultaneous acquisition of different MS–MS experiments increasing the information obtained in a single run [64], and the development of novel bioinformatic tools that ensure complex data mining [65].

Among the different direct DBS techniques discussed above, on-line DBS and SSSP (i.e. the Camag interface) combined with LC separation capacity, seem to be

particularly convenient for identification of compounds in complex biomatrices. In our laboratory, we recently developed a TS strategy that enabled the identification and semi-quantification of 90 selected pharmaceuticals and metabolites by automated on-line DBS LC–MS–MS, with a run time of 10 min. Representative MRM transitions and the estimated LOQ are listed in Table 2 and the complete list is given in Electronic Supplementary Material Table S1. All of the compounds of interest were tested using automated on-line DBS with RP-LC separation before MS–MS detection in ESI mode.

### Preclinical study

With regard to the recent implementation of the DBS sampling process in the pharmaceutical industry, it is apparent that DBS are becoming increasingly important in preclinical studies of drug metabolism and pharmacokinetics (DMPK) [66]. In the early stages, drug candidates are tested for ADME properties using small animal models, in which amounts of biological material are often restricted [16]. Thus, if a conventional sampling procedure is used (plasma), one



**Fig. 5** Typical workflow of a clinical study performed with the DBS sampling process followed by on-line LC–MS–MS analysis (A). For this trial, 50 mg flurbiprofen was orally administered to human volunteers and DBS samples were collected over eight hours. Representative log

concentration–time profiles for flurbiprofen (triangles) and the OH-flurbiprofen metabolite (squares), and the associated MRM chromatograms at two hours are presented (B)

animal must be sacrificed for each experiment or time-point in a kinetic study. With DBS sampling, multiple experiments or time-points can be performed on the same animal, leading to significant benefits in the 3Rs (reduction, refinement, and replacement) in animal use [67]. Moreover, direct DBS techniques now enable increased throughput of DBS analysis; this is particularly appropriate in the pharmaceutical industry, in which large numbers of drug candidates are tested. For example, Roche laboratories have successfully applied on-line DBS using SCAP technology to a study of the pharmacokinetic of oseltamivir on rats [43].

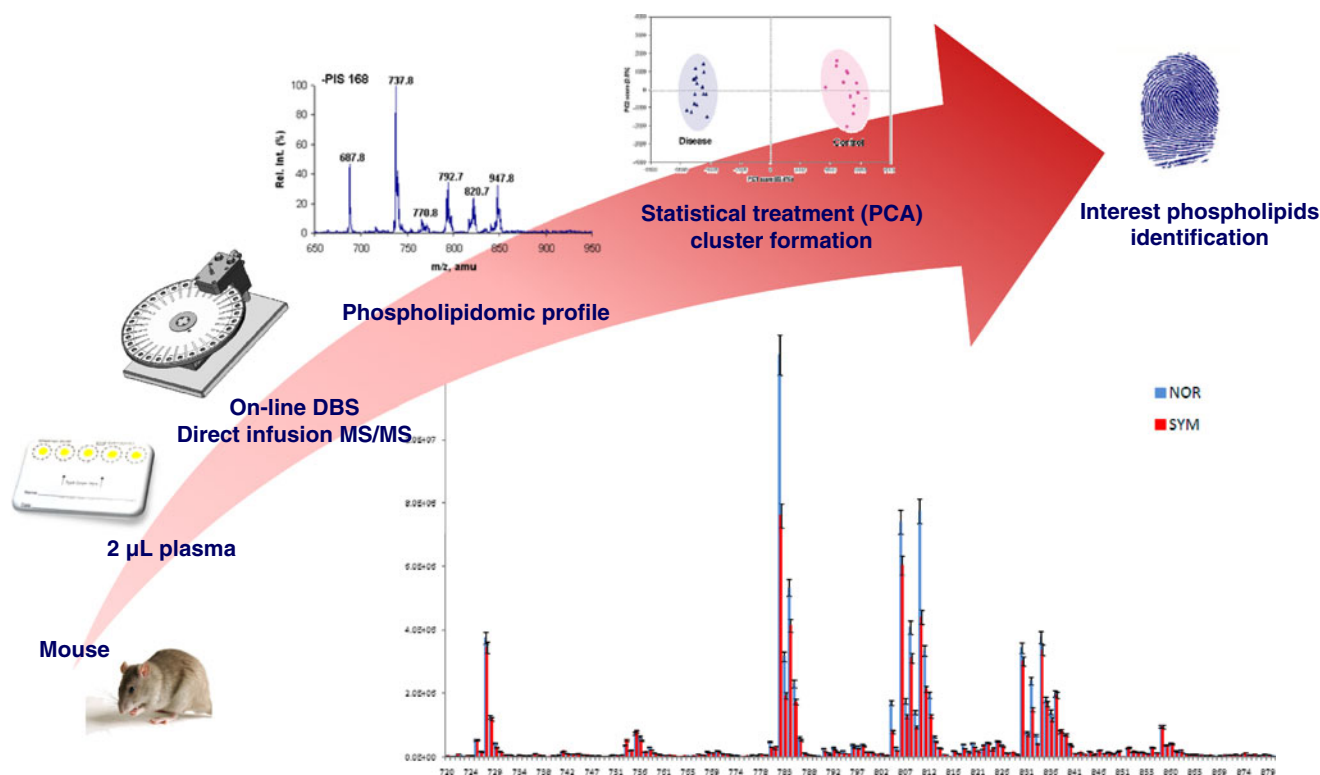
Even though very few papers have been published focusing on drug metabolism studies based on DBS [68], Thomas et al. demonstrated the possibility of highlighting phase I and II metabolites of buprenorphine in rats after intraperitoneal injection. In these experiments, buprenorphine, norbuprenorphine, and their respective glucuronides were simultaneously monitored from 5- $\mu$ L DBS samples [15].

### Clinical study and therapeutic drug monitoring

If the ethical benefit of using DBS is clearly established for animal use in preclinical studies, this less invasive sampling mode can be easily applied to human clinical studies or

TDM investigations. As reviewed by Li and Tse, DBS–LC–MS–MS has been increasingly used in this area for analysis of a wide range of drug molecules, including antiretroviral, immunosuppressant, antiepileptic, or antimalarial drugs [19]. Moreover, Glaxo Smith Kline regards DBS sampling as the preferred sampling method for orally dosed compounds in Phase I clinical trials [69]. Although all the DBS methods discussed in this section are based on off-line extraction with subsequent LC–MS–MS analysis, direct strategies, discussed in this review, will certainly promote DBS analysis in clinical laboratories. For example, Roche has successfully applied the Camag TLC–MS interface to phase I determination of drug candidates in human DBS samples [44].

Furthermore, this kind of strategy may be of important benefit to clinical pharmacology in terms of ease of use for medical staff and patient comfort. On the basis of these considerations, an automated on-line DBS method for pharmacokinetic study of flurbiprofen and its main metabolite 4-hydroxyflurbiprofen in human volunteers was developed at the University Hospitals of Geneva to assess the activity of cytochrome P450 2C9 (Fig. 5). The system was successfully tested on over 500 real samples, demonstrating the good competitiveness of the process compared with conventional plasma LC–MS–MS analysis [39].



**Fig. 6** High-throughput phospholipidomic fingerprinting for evaluation of disease biomarkers. For this purpose, 2  $\mu$ L mouse plasma previously spotted on filter paper was analyzed by MS–MS after direct infusion using an on-line DBS process. Multiple MS–MS

experiments (neutral loss and precursor ion scans) were acquired for simultaneous measurement of phospholipidic classes. Complex data mining was performed by use of statistical tools to ensure sample classification and to highlight the discriminated species

## Identification of biomarkers

The discovery of novel biomarkers is of major importance for diagnosis of disease states, determination of prognosis, and therapeutic response monitoring [70]. Over the past decade, LC–MS–MS has become the key analytical technique in the investigation of biomarkers by proteomic and metabolomic approaches [71]. Regarding implementation of DBS in LC–MS–MS workflow (as discussed previously), direct DBS analysis may be a novel means of supporting the different stages in the biomarker pipeline from discovery to the validation steps [72]. Preliminary tests have demonstrated the feasibility of the analyses of the enzymes and peptides spotted on filter paper using the paper spray process [58]. Furthermore, Thomas et al. developed an on-line DBS MS–MS strategy for high-throughput fingerprinting of phospholipidome in atherosclerotic mice from 2 µL dried plasma that was desorbed directly into the MS instrument without pre-treatment or LC separation (Fig. 6) [40]. Based on multiple MS–MS experiments and data mining, this study has enabled sample classification between atherosclerotic and control mice and highlighted several species that were significantly up and down-regulated as a consequence of that pathology.

## Conclusion

Over the past decade, DBS sampling has emerged as a powerful alternative approach for clinical and pharmaceutical analysis compared with the conventional venipuncture procedure. The filter paper affords numerous advantages that range from better shipment/storage logistics to a less invasive and more ethical procedure. However, the lack of automation, necessitating manual off-line extraction, could make this procedure unsuitable for routine laboratories. The recent developments highlighted in this review enable direct MS–MS analysis of DBS samples (with or without previous LC separation) and offer competitive solutions in terms of throughput and sensitivity compared with the plasma procedure. Furthermore, the marketing of automatic methods indicates that some direct DBS methods are now mature and can be easily implemented in analytical laboratories.

**Conflict of interest/disclosure** The authors declare no conflicts of interest regarding this manuscript.

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