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#### Review

## Coupling continuous separation techniques to capillary electrophoresis

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#### **Abstract**

One of the weak points of capillary electrophoresis is the need to implement rigorously sample pretreatment because its great impact on the quality of the qualitative and quantitative results provided. One of the approaches to solve this problem is through the symbiosis of automatic continuous flow systems (CFSs) and capillary electrophoresis (CE). In this review a systematic approach to CFS–CE coupling is presented and discussed. The design of the corresponding interface depends on three factors, namely: (a) the characteristics of the CFS involved which can be non-chromatographic and chromatographic; (b) the type of CE equipment: laboratory-made or commercially available; and (c) the type of connection which can be in-line (on-capillary), on-line or mixed off/on-line. These are the basic criteria to qualify the hyphenation of CFS (solid-phase extraction, dialysis, gas diffusion, evaporation, direct leaching) with CE described so far and applied to determine a variety of analytes in many different types of samples. A critical discussion allows one to demonstrate that this symbiosis is an important topic in research and development, besides separation and detection, to consolidate CE as a routine analytical tool. © 2001 Elsevier Science B.V. All rights reserved.

Keywords: Sample handling; Leaching; Dialysis; Evaporation; Extraction methods; Membranes; Isotachophoresis; Reviews

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#### 1. Introduction

The analysis of real samples by capillary electrophoresis (CE) requires efficient sample preparation procedures to remove interfering solutes, (in)organic salts and particulate matter. Sample preparation systems reported in the literature are based on chromatographic, electrophoretic, membrane-based procedures, solid-phase extraction (SPE), supercritical fluid extraction (SFE) among others techniques. The combination of automated sample preparation and CE is especially useful if complex samples have to be analysed and helps to improve both selectivity and sensitivity. In this review, the different modes of sample pretreatment will be discussed and an overview of the potential of these procedures will be given.

The number of routine applications of real samples using CE is limited because there still are several problems that have to be solved. No doubt the main problem is that analyte detectability expressed in concentrations units, generally is rather poor because of the low volume loadability of the capillary. Another problem is the presence of particulate matter, which can easily clog the CE system [1]. The emphasis of this review will be on those sample preparation procedures that can be used for the quantitative determination of analytes in real samples.

This review is not intended to provide readers a detailed description of the published papers; rather, it has tried to summarise in different tables more than 450 references which were classified according to the coupling mode, the type of the continuous flow system (CFS) used, the electrophoretic mode, type of sample and analytes. Moreover, tables will help to organise the subject and emphasise the primary events. Only documents published since 1980 up to beginning year 2000 will be considered in this review. The rapid growth in the number of publications for CE has forced this review to limit the number of papers cited and to concentrate primarily on those documents which the emphasis is on the coupling of CFSs with CE.

#### 2. Functions of continuous flow systems

The use of low-pressure continuous flow systems

can be viewed as one of the first reliable steps towards real performance improvement in analytical methods through the automation, miniaturisation and simplification of the preliminary operations of analytical methodologies. These systems allow the implementation of one or several (simultaneous or sequential) chemical reactions, as well as reliable separation techniques for increasing sensitivity and selectivity, calibration procedures, etc. Also, CFSs have recently proved to be useful tools for developing rapid response analytical systems such as sensors and screening devices [2].

The favourable influence of CFSs on analytical properties [3] is undeniable. In fact they provide a highly useful means for improving: (a) the productivity of analytical laboratories through increased throughput, personnel safety and comfort, and reduced costs, and (b) the accuracy of the analytical results through increased precision, sensitivity and selectivity. In summary, CFSs can be included among the most outstanding advances in chemical measurement processes.

There are two main ways of combining continuous flow systems and instruments. In the simpler one, the flow system is connected via an appropriate interface to an instrument that is the exclusive source of analytical information. More powerful is the coupling of a flow system furnished with a non-destructive detector to an instrument of a high information level allowing physical or physicochemical discrimination of analytes and providing reliable qualitative and/or quantitative information. The global information furnished by the detector in the continuous flow system can be used for a variety of purposes [2].

### 3. Continuous flow system-capillary electrophoresis coupling modes

CE is a highly flexible and efficient analytical separation technique that has become a serious competitor for gas, liquid and supercritical column chromatographies. Nevertheless, its intrinsic features make direct sample introduction almost impossible. Sample pretreatment is crucial in order to obtain reliable analytical results. Thus, CFSs linking gaseous, liquid gaseous, liquid or solid samples to the CE system are of great practical importance because they

can automatically implement such operations as dissolution, leaching, filtration, derivatisation, matrix isolation, analyte concentration, solvent exchange and so on. Continuous separation techniques such as SPE, dialysis, etc., play major roles in this context.

Although CFSs and CE are of hydrodynamic nature, there are many technical aspects that make them theoretically incompatible. Nevertheless, several types of interfaces have been described for coupling CFS–CE and it were summarised by our group [2].

Combinations of a CFS and CE equipment can be characterised by the degree of integration between these two units. Four levels of coupling can be distinguished: off-line, at-line, on-line and in-line. The sample preparation modules can be coupled with CE either off-line (manual), at-line (robotic interface) on-line (coupling via a transfer line) or in-line (complete integration between sample preparation and separation system).

#### 4. Leaching coupled to capillary electrophoresis

A few determinations of analytes in solid samples have been find in the literature. In Table 1, five references are summarised.

An automatic system for stepwise treatment of solid samples and application to pollution evaluation by measuring ion lixiviation rates in lichens by capillary zone electrophoresis (CZE) was developed by Arhoun et al. [4]. Lichen (0.4 g) was immersed in 10 ml water in an extraction tube and sonicated in a bath at 30°C for 20 min. The supernatant was filtered and hydrodynamically injected. The cation lixiviation rates are related to their exposure to pollution.

Recently, a CE method was developed for the simultaneous determination of a number of major ingredients of green tea. Analysis was carried out after treatment (extraction, filtration and dilution) of the samples in a flow system which was coupled to a commercial CE equipment via a programmable arm [5].

Another example of leaching coupled to CE was reported by Aguilar et al. [6]. Capillary electrophoretic determination of cyanide leaching solutions from automobile catalytic converters was carried out. A CE method was developed for determining Fe(II)-, Cu(I)-, Ni(II)-, Pd(II)- and Pt(II)-cyano complexes and nitrite in the leaching solutions.

#### 5. Dialysis coupled to capillary electrophoresis

Dialysis is normally used to remove particulate material and can be coupled with a wide variety of separation techniques. Microdialysis has emerged as a powerful tool for monitoring the extracellular environment of a variety of organs, tissues and bodily fluids in vivo. Microdialysis is powerful because it can be coupled with a variety of analytical methods; therefore, it can be used to simultaneously monitor a large variety of endogenous and exogenous compounds. Techniques that have been coupled to the dialysis probe include LC, immunoassay, mass spectrometry (MS) and CE, an example is shown in Fig. 1 [7]. The usefulness of the dialysis for the pretreatment of the samples before its introduction into the CE equipments is shown with more than 45 references in Table 2.

A flow-gated, on-line interface between a microdialysis sampling probe and CE with UV de-

Table 1 Coupling continuous leaching systems to capillary electrophoresis

| Coupl | ing m | ode |        | CFS  | Electrophoretic | Sample             | Analytes       | Remarks  | Ref. |
|-------|-------|-----|--------|------|-----------------|--------------------|----------------|--|------|
| OFF   | IN    | ON  | OFF/ON | NC C | mode            |                    |                |  |      |
| x     |       |     |        | X    | CZE             | Lichen             | Cations        | Determination of lixiviation rates of K <sup>+</sup> , Na <sup>+</sup> , Ca <sup>2+</sup> and Mg <sup>2+</sup> from the lichen | [4]  |
|       |       | X   |        | X    | CE              | Tea                | Polyphenols    | Extraction of analytes with water from a solid sample (tea)  | [5]  |
| X     |       |     |        | X    | CE              | Car exhaust gases  | PGMs           | Analytes recovered from automobile catalytic converters by a leaching process  | [6]  |
| X     |       |     |        | X    | CE              | Metal cyanide      | Au, Ag, Cu, Ni | Analytes were determined in samples from the leaching of Au numeral and sand   | [58] |
| X     |       |     |        | X    | CE              | Aerosols/rainwater | Iron (II)      | Sample was prepared for CE by leaching with 0.2 ml of HCl and 2.0 ml of water  | [59] |

CFS: Continuous flow system; NC: no chromatography; C: chromatography; CZE: capillary zone electrophoresis; PGMs: platinum group metals

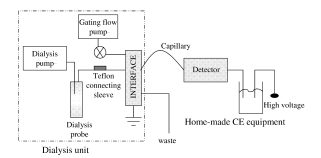


Fig. 1. On-line interface between microdialysis and capillary zone electrophoresis (adapted from Ref. [7]).

tection was characterised and applied. The data presented by Lada et al. [7] demonstrated the utility of the interface for allowing low perfusion flow-rates while allowing temporal resolution of 65–85 s.

An on-line coupling of microdialysis sampling to CE was described by Hogan et al. [8]. The heart of the on-line system is the interface between the microdialysis apparatus and the CE system. The interface must isolate the experimental animal from the high potential of the CE experiment and convert the microliter per minute microdialysis flow into discrete nanoliter volume samples for the CE while not adding to the system's dispersion. This was accomplished by using a commercially available rotary microinjection valve for LC with an injection apparatus designed in the laboratory.

In 1997, an automated method for high temporal resolution monitoring of the neurotransmitters glutamate and aspartate in vivo using CE with laser-induced fluorescence (LIF) detection was developed [9]. Microdialysis probes placed in the striatum of anaesthetised rats were coupled on-line with the CE system by an automated flow-gated interface. Flexible loop microdialysis probes (ESA, Bedford, MA, USA) made from cellulose fibres ( $M_r$  6000 cutoff) with 450  $\mu$ m O.D. tip diameters and 2 mm tip lengths were used for all sampling experiments. The dialysis membrane had a 210  $\mu$ m I.D. for an internal volume of 0.12  $\mu$ l.

In the same year, a CE-electrospray ionisation (ESI) MS interface, based on an electric circuit across a microdialysis membrane surrounding a short capillary segment closely connected to the separation capillary terminus was demonstrated by Severs and Smith [10] to be sensitive, efficient, and rugged.

Kuban and Karlberg [11] described a CE application using a common electrolyte for separation of the most common small inorganic/organic anions and cations. The system uses one capillary and just one detector placed in the centre of the capillary. The technique has successfully been applied to the simultaneous determination of anions and cations in natural water samples. Milk and mud samples pretreated by dialysis have also been analysed. On-line dialysis performed in a flow injection analysis (FIA) system was integrated with a CE system via a specially designed interface, developed by the same authors [12]. Samples were continuously pumped into a dialysis unit and the outgoing acceptor stream containing the analytes is allowed to fill a rotary injector in the FIA part of the system. A discrete, representative volume of the acceptor stream is injected into an electrolyte stream, which continuously passes through the FIA-CE interface into which the end of a capillary has been inserted.

A fully automated method is presented for the determination of acidic drugs in urine and serum using on-line dialysis—SPE—CE [13]. This system is fully automated and 250–500 analyses can be carried out without any maintenance or exchange parts.

In 1999, a rapid determination of aspartate enantiomers in tissue samples by microdialysis coupled on-line with CE was developed by Thompson et al. [14]. The microdialysis probe was inserted into a homogenised tissue sample, which allowed generation of a continuous, sample stream that was filtered and deproteinated. Values of D- and L-aspartate in different tissues agreed well with those obtained by a high-performance liquid chromatography (HPLC) procedure that required protein precipitation, centrifugation, and extraction. The speed and compatibility with automation of the microdialysis—CE method may make it a general approach for a variety of applications involving high-throughput analysis or sensorlike operation.

Finally Crowder et al. [15] developed a method for the determination of phosphoamino acids with indirect photometric detection. The samples were hydrolysed in HCl, aliquots of the cooled hydrolysis solution were dialysed for 3 h in  $M_{\rm r}$  100 cutoff Spectra/Por Biotech cellulose ester dialysis tubing. A portion of this solution was injected into the CE instrument.

Table 2 Coupling continuous dialysis systems to capillary electrophoresis

| Coupl | ling m | ode |        | CFS |   | Electrophoretic | Sample              | Analyte                              | Remarks   | Re     |
|-------|--------|-----|--------|-----|---|-----------------|---------------------|--------------------------------------|---|--------|
| OFF   | IN     | ON  | OFF/ON | NC  | С | mode            |                     |                                      |   |        |
|       |        | X   |        | X   |   | CE              | Rat brain           | Ascorbic acid                        | Microdialysis sampling probe                          | [7     |
|       |        | X   |        | X   |   | CE              | Rats                | Pharmacokinetics products            | Cellulose fibre dialysis                              | [8     |
|       |        | X   |        | X   |   | CE              | Rats                | Glutamate, aspartate                 | Microdialysis probes                                  | [9     |
|       |        | X   |        | X   |   | CE              | _                   | Proteins                             | Interface based on microdialysis membrane             | [1     |
|       |        |     |        | X   |   | CE              | Water, milk, mud    | Cations and anions                   | CE laboratory-made                                    | [:     |
|       |        | X   |        | X   |   | CE              | Water, snow, mud,   | Small anions                         | Microdialysis CE laboratory-made                      | [:     |
|       |        | X   |        | X   |   | CE              | Serum, urine        | Acidic drugs                         | Cellulose acetate membrane                            | [:     |
|       |        | X   |        | X   |   | CE              | Tissue              | Aspartate enantiomers                | Microdialysis coupled with CE                         | [      |
|       |        |     |        | X   |   | CE              | _                   | Phosphoamino acids                   | -   | ſ      |
|       |        |     |        | X   |   | CZE             | Royal jelly         | Proteins                             | _   | [      |
|       |        |     |        | X   |   | MEKC            | Polymers            | Alkylphenyl ketones, alkyl benzoates | Products were purified by dialysis                    | [      |
|       |        | X   |        | x   |   | IEF             | _                   | Proteins                             | Microdialysis sleeve tubing                           | [(     |
|       |        | -   |        | X   |   | CE              | Plasma              | Drug-protein                         | Review  | [      |
|       |        |     |        | X   |   | CE              | Tissues, organs     | Pharmacokinetic                      | Review  | [0     |
|       |        | X   |        | X   |   | CE              | Live animal         | Nicotine                             | Rotary switching valve                                | [      |
|       |        | X   |        | X   |   | CE              | Cerebrospinal fluid | Gabapentin (I)                       | Microdialysis probe                                   | [      |
|       |        | X   |        | X   |   | CE              | Blood serum         | Sulfonamides                         | Dialysis-SPE device                                   | 1      |
|       |        | X   |        | Λ   | x | CE              | Diood scruiii       | Collagen crosslinks                  | Three detection modes                                 | 1      |
|       |        | А   |        | v   | λ | CE              | _                   | Ethoxylated polymers                 | Cellulose ester dialysis tube                         | ا<br>ا |
|       |        |     |        | X   |   | IEF             | Dhamaluta 5 0       |                                      |   |        |
|       |        | X   |        | X   |   |                 | Pharmalyte 5-8      | Carbonic anhydrase                   | Plexiglas microdialysis chamber                       |        |
|       |        |     |        | X   |   | CE              | Serum               | Oligonucleotides                     | Membrane filters (cellulose)                          |        |
|       |        |     |        | X   |   | CE              | Blood plasma        | Oligonucleotides                     | Drop dialysis   | - 1    |
|       |        | X   |        | X   |   | CE              | -                   | Myoglobin, C anhydr.                 | Piece of dialysis tubing                              | [      |
|       |        |     |        | X   |   | CZE             | Rat brain           | Glutamate                            | -   | [      |
|       |        | X   |        | X   |   | CE              | -                   | Benzenesulfonamides                  | Sleeve of polysulfone dialysis                        | - 1    |
|       |        |     |        |     |   | CE              | -                   | _                                    | Review  |        |
|       |        |     |        | X   |   | CE              | PCR products        | Nucleic acids                        | Membrane dialysis                                     | -      |
|       |        |     |        | X   |   | CE              | Smoked fish         | Clostridium botulinum                | Membrane dialysis                                     | -      |
|       |        |     |        | X   |   | CGE             | DNA                 | Clostridium botulinum                | Membrane dialysis                                     | -      |
|       |        |     |        | X   |   | CGE             | Human plasma        | Phosphorothioate oligonucleotides    | Millipore VS membrane                                 |        |
|       |        |     |        | X   |   | CE              | Living rat brain    | γ-Aminobutyric acid                  | In vivo microdialysis                                 |        |
|       |        |     |        | X   |   | CE              | Blood               | $\alpha$ -Difluoromethylornithine    | Microdialysis   |        |
|       |        |     |        | X   |   | CE              | Proteins            | -                                    | Dialysis memebrane                                    | - 1    |
|       |        | X   |        | X   |   | IEF             | Proteins            | -                                    | Hollow dialysis fibre                                 | - 1    |
|       |        |     |        | X   |   | CE              | Blood serum         | Organic anions                       | Dialysis fluids                                       |        |
|       |        |     |        | X   |   | CE              | Animals             | Aspartate and glutamate              | Microdialysis sampling                                | - 1    |
|       |        |     |        | X   |   | CE              | Peas                | Soyasaponin I                        | _   | - 1    |
|       |        |     |        | X   |   | CE              | DNA                 | Polymerase chain reaction            | -   | -      |
|       |        |     |        | X   |   | ITP-CZE         | Kidney cells        | Antithrombin III                     | Samples purified by dialysis                          |        |
|       |        |     |        | X   |   | CE              | Brain rats          | Glutamic acid                        | Microdialysis probe sampling                          |        |
|       |        |     |        | x   |   | CE              | Brain               | Glutamate I                          | Microdialysis coupled to CE                           | [      |
|       |        |     |        | x   |   | CE              | Rats                | Aspartate, glutamate and alanine     | Microdialysis   | [      |
|       |        |     |        | x   |   | CE              | Rats                | L-dopa                               | In vivo microdialysis                                 |        |
|       |        |     |        | x   |   | CGE-ITP         | Roasted coffee      | Total titratable acid                | -   | [      |
|       |        |     |        | X   |   | CE              | Brain               | Glutamate                            | Microdialysis and CE                                  | [      |
|       |        | X   |        | X   |   | CE              | Rat caudate nucleus | Primary amines                       | Microdialysis coupled by a flow-gated interface to CE |        |

CFS: Continuous flow system; NC: no chromatography; C: chromatography; CZE: capillary zone electrophoresis; CE: capillary electrophoresis; MEKC: micellar electrokinetic chromatography; IEF: capillary isoelectric focusing; CGE: capillary gel electrophoresis; IEF: capillary isoelectric focusing; PCR: polymerase chain reaction; SPE: solid-phase extraction; ITP: capillary isotacophoresis.

### 6. Evaporation coupled to capillary electrophoresis

In the classical analyte extraction protocols from real samples, a great number of organic solvent are used. The analytes dissolved in theses solvents are in almost all cases incompatible with the buffer used in the CE method. Due to this reason is compulsory the evaporation of these solvents and the redisolution in solvent compatible with the CE methodology. Some examples are shown in Table 3.

The determination of methotrexate and its major metabolite, 7-hydroxymethotrexate using CE and LIF detection was carried out by Roach et al. [16]. Serum was mixed with  $0.2\ M$  sodium acetate (pH 5.1) and the mixture was cleaned up on a Sep-Pak  $C_{18}$  cartridge with elution with methanol. The eluate was evaporated to dryness and the residue was dissolved in  $1\ M\ 2$ -(N-morpholino)ethanesulfonic

acid (I) and 0.35%  $\rm KMnO_4$  was added to oxidise the analytes.

Micellar electrokinetic capillary electrophoresis (MEKC) provides rapid and efficient separation and detection of organic gunshot and explosive constituents [17]. Samples for MEKC analysis were obtained by ultrasonic agitation of the swab in 500 μl of ethanol. The ethanol was evaporated down to a 2–3 μl under a stream of nitrogen and then diluted with 50 μl of the running buffer.

Zhang and Hjerten [18] developed a micro method for concentration and desalting utilising a hollow fibre with special reference to CE. Aqueous sample, e.g., K<sub>2</sub>CrO<sub>4</sub> or protein, respectively, was introduced into a hollow fibre and concentrated by spontaneous or forced evaporation of water through the fibre wall or by Donnan transport of water into a polymer solution surrounding the fibre. The concentrated solutions were diluted if required and subjected to CE (procedures described). The possibility of using

Table 3 Coupling continuous evaporation systems to capillary electrophoresis

| Coupl | ing m | ode |        | CFS |   | Electrophoretic | Sample                     | Analyte                                   | Remarks                                 | Ref.  |
|-------|-------|-----|--------|-----|---|-----------------|----------------------------|---|---|-------|
| OFF   | IN    | ON  | OFF/ON | NC  | С | mode            |                            |   |   |       |
| Х     |       |     |        | Х   |   | CZE             | Serum/blood                | Methotrexate and its 7-hydroxy metabolite | Eluate was evaporated to dryness        | [16]  |
| X     |       |     |        | X   |   | MEKC            | Swabs                      | Organic gunshot                           | Ethanol evaporated under a stream of N2 | [17]  |
|       |       | X   |        | X   |   | CE              | Low or high molecular mass | -   | Hollow fibre, forced evaporation        | [18]  |
| X     |       |     |        | X   |   | CE              | Urine                      | Tramadol and its main metabolites         | Residue evaporation                     | [97]  |
| X     |       |     |        | X   |   | CE              | Human plasma               | Doxorubicin (I)                           | Evaporation of organic phase            | [98]  |
| X     |       |     |        | X   |   | CZE             | Corn                       | Tosylated polyamines                      | Alkaline solution evaporated            | [99]  |
| X     |       |     |        | X   |   | CE              | _                          | Dansyl-D-L-glycine/valine                 | Solvent evaporation                     | [100] |
| X     |       |     |        | X   |   | CZE             | Magnolia officinalis       | Magnolol and honokiol                     | Evaporation of ethanol                  | [101] |
| X     |       |     |        |     | X | GPC             | Vesicle                    | Dextrans in lipid                         | Reversed-phase evaporation              | [102] |
| X     |       |     |        | X   |   | CE              | Organic solvents           | Acidic species                            | Evaporation of volatile solvents        | [103] |
| X     |       |     |        | X   |   | CE              | _                          | -   | Evaporation of buffer solvent           | [104] |
| X     |       |     |        | X   |   | MEKC            | Citrus seeds               | Limonoid glucosides                       | Soxhlet extracted                       | [105] |
| X     |       |     |        | X   |   | CZE             | Butter                     | Free fatty acids                          | _                                       | [106] |
| X     |       |     |        | X   |   | MEKC            | Soil samples               | PAHs                                      | Solvent evaporation                     | [107] |
| X     |       |     |        | X   |   | CE              | Glycoproteins              | Oligosaccharides                          | Evaporation of excess reagent           | [108] |
| X     |       |     |        | X   |   | CZE             | Serum                      | Polyamines                                | Evaporation of acetone                  | [109] |
| X     |       |     |        | X   |   | MEKC            | Plasma, human milk         | Thiocyanate, iodide, nitrate and nitrite  | Solvent evaporation                     | [110] |
| X     |       |     |        | X   |   | CE              | Glycoproteins              | Oligosaccharides                          | Solvent evaporation                     | [111] |
| X     |       |     |        | X   |   | MEKC            | Seeds and plants           | Glucosinolates                            | Evaporation from samples                | [112] |
| X     |       |     |        | X   |   | CZE             | _                          | Amino acids                               | _                                       | [113] |

CFS: Continuous flow system; NC: no chromatography; C: chromatography; CZE: capillary zone electrophoresis; CE: capillary electrophoresis; MEKC: micellar electrokinetic chromatography; GPC: high-performance gel exclusion chromatography; PAHs: polycyclic aromatic hydrocarbons.

the method for on-line preconcentration for capillary electrophoresis is discussed.

### 7. Supercritical fluid extraction coupled to capillary electrophoresis

The application of multidimensional chromatography to the analysis of complex matrices helps to minimise sample pretreatment steps. Still, when the matrix to be analysed is not totally soluble in a particular solvent, such as for example plant tissue, a preliminary step is necessary to obtain a solution suitable for subsequent introduction into the preliminary separation stage. Supercritical fluids offer potential advantages over liquid solvents to meet the sample preparation requirements. SFE was used in conjunction with thin-layer chromatography and conventional LC. The on-line coupling of SFE to gas chromatography (GC) is experiencing rapid growth and will continue to be studied as a simplified method of sample preparation and analysis [19]. To our knowledge only 12 articles used SFE to extract the analytes which were separated by CE (see Table 4).

In the last 2 years, our team has worked on the coupling SFE-CE. An automatic method for the

determination of phenols in liquid samples (river water and human urine) using co-electrosmotic CE coupled via a laboratory-mechanical arm to a supercritical fluid extractor, was developed and validated. Samples were preconcentrated onto  $C_{18}$  sorbent and carefully transferred to extraction cartridges for extraction with supercritical  $CO_2$ . The analytes were collected in a diol-trap and eluted with methanol, which is synchronically fed to the CE vial by the mechanic arm, controlled via an electronic interface [20].

In 1996 solid-liquid extraction and SFE carried out the determination of carbamate residues in tobacco samples. The results were analysed by CE and the results compared, demonstrating the advantages of using SFE to reduce time, expense, hazardous wastes and enhance extraction power [21].

In the same year, a separation method using cyclodextrin-modified CE has been developed for analysis of the polycyclic aromatic hydrocarbons (PAHs) in contaminated soils by Brown et al. [22]. Contaminated soil was extracted using  ${\rm CO_2}$  supercritical fluid. The diluted extracted was analysed using fluorescence detection.

Finally, Wang and Chang [23] described an offline SFE–CE procedure for the determination of four parabens in cosmetics samples.

| Table 4                           |                          |                                |
|-----------------------------------|--------------------------|--------------------------------|
| Coupling continuous supercritical | fluid extraction systems | s to capillary electrophoresis |

| Couplin | ng mode | ;  |        | CFS |   | Electrophoretic | Sample              | Analyte                     | Remarks                                | Ref.  |
|---------|---------|----|--------|-----|---|-----------------|---------------------|-----------------------------|--|-------|
| OFF     | IN      | ON | OFF/ON | NC  | С | mode            |                     |                             |  |       |
|         |         | X  |        | X   |   | CE              | Phenols             | Water and urine             | Laboratory mechanical arm              | [20]  |
| X       |         |    |        | X   |   | CZE             | Residues in tobacco | Carbamates                  | Supercritical CO <sub>2</sub> modified | [21]  |
| X       |         |    |        | X   |   | CE              | Contaminated soils  | PAHs                        | Cyclodextrin-modified                  | [22]  |
| X       |         |    |        | X   |   | CZE             | Cosmetic products   | Parabens                    | Supercritical CO <sub>2</sub> modified | [23]  |
| X       |         |    |        | X   |   | CE              | _                   | Ethoxylated polymers        | Excess reagent was removed by SFE      | [69]  |
| X       |         |    |        | X   |   | MEKC            | Blood stains        | Smokeless powder residues   | _                                      | [114] |
| X       |         |    |        | X   |   | MEKC            | Food                | Vitamins                    | _                                      | [115] |
| X       |         |    |        | X   |   | CE              | Water               | Carboxylic acids            | Supercritical water                    | [116] |
|         |         |    |        |     |   | CE              | Environmental       | -                           | Review (solid matrices)                | [117] |
| X       |         |    |        | X   |   | CZE             | Water               | Phloxine B and uranine (II) | SFE was used for spiked water samples  | [118] |
| X       |         |    |        | X   |   | CE              | Stevia rebaudiana   | Steviol glycosides          | Subcritical fluid extraction using CO2 | [119] |
| x       |         |    |        | X   |   | CE              | Antioxidative       | Plant beverages             | Leaf extracts were prepared by SFE     | [120] |

CFS: Continuous flow system; NC: no chromatography; C: chromatography; CZE: capillary zone electrophoresis; CE: capillary electrophoresis; MEKC: micellar electrokinetic chromatography; PAHs: polycyclic aromatic hydrocarbons; SFE: supercritical fluid extraction.

### 8. Solid-phase extraction coupled to capillary electrophoresis

SPE can be used to simultaneously enrich the trace analytes and remove potentially interfering compounds. As can be seen in Table 5, it can be combined with CE in several ways that vary from off-line to in-line approaches. To circumvent poor CE limit of detection off-line sample pretreatment and analyte concentration could be achieved. However, if possible, this should be avoided for dilute analyte solutions since losses to exposed surfaces (e.g., walls of Eppendorf tubes, pipette tips, solid extraction phases, etc.) can be substantial. To avoid these problems, minimal sample handling is advisable. This can be achieved using an analyte concentrator [24,25] on-line with the CE capillary. These devices usually consist of an adsorptive phase at the inlet of the CE capillary and serve to enrich trace levels of analytes, as well as allow on-line sample clean up prior to component separation by CE.

In 1997, a sensitive method for the determination of PAHs by solid-phase microextraction (SPME) coupled with cyclodextrin-modified CE using UV detection has been developed [26]. A glass fibre was prepared and used for absorbing 16 US Environmental Protection Agency (EPA) priority PAHs from diluted samples until equilibrium was reached. After the glass fibre was connected to a separation capillary via an adapter, the absorbed analytes were directly released into the CE buffer stream. Very satisfactory reproducibility with respect to migration time and peak area was obtained for repetitions using the same separation capillary and adapter, where only the extraction fibre was discarded after each analysis.

One year later, an on-column interface, coupling the SPME sampling technique with CE was constructed [27]. This interface facilitates the direct insertion of a thin silica fibre into the inlet end of a separation capillary and, therefore, the zero-dead volume connection requirement for hyphenation between SPME and CE was fully realised. The performance of the interface was evaluated by SPME–CE analysis of the priority pollutant phenols using a laboratory-made 40 µm O.D. poly(acrylate) (PA)-coated silica fibre connected to a 75 µm I.D.

separation capillary. The results clearly demonstrated that the interface was effective for on-line coupling SPME with CE.

In the same year, a separation of chiral biodegradation intermediates of linear alkylbenzenesulfonates by CE was carried out by Kanz et al. [28]. After the enrichment on graphitised carbon black material, the extracts were analysed by HPLC and CE.

In our laboratory, a CZE method was developed for the determination of heterocyclic aromatic amines in meet and fish sample. SPE (the Gross methodology) was tested for isolating the amines [29]. The same authors [30] developed also a method for the determination of chlorophenols (CPs) in human urine by using MEKC coupled via a mechanic arm to an on-line automatic clean up and preconcentration unit for urine samples. The coupling of both systems allows the expeditious, reproducible, sensitive, and inexpensive determination of CPs in human urine with acceptable precision and accuracy. In the same laboratory, different flow injection (FI) systems furnished with different minicolumns (C<sub>18</sub> and Chelex-1000) were used to clean up the samples and preconcentrate analytes. The analytes eluted from the solid phases were driven from the FI systems to the autosampler of the CE equipment by a programmable arm [31-34], an example is shown in Fig. 2. This interface can be considered as a general approach to make discrete analytical equipment (which they commonly use sample turntables) compatible with the continuous flow system.

The FI-SPE-CZE system described by Chen and Fang [35] was shown to be applicable to the automated preconcentration, separation and determination of trace amounts of drug constituents in blood plasma, as demonstrated by the quantitation of pseudophedrine in plasma down to the 50 µg l<sup>-1</sup> level. The FI-SPE system not only concentrated the analyte but also modified the sample medium to create the low conductivity required for further sensitivity enhancement by electro-stacking.

On-line ion-exchange preconcentration, performed in a FIA system, has been integrated with laboratorymade CE equipment via a specially designed interface. A sensitive and selective method for the determination of nitrite, nitrate, bromide and iodide

Table 5 Coupling continuous solid-phase extraction systems to capillary electrophoresis

| Coup | ling n | node |        | CFS | Electrop   | phoretic Sample          | Analytes                                      | Remarks   | Ref |
|------|--------|------|--------|-----|------------|--------------------------|---|---|-----|
| OFF  | IN     | ON   | OFF/ON | NC  | C mode     |                          |   |   |     |
| X    |        |      |        | х   | CE         | Serum and urine          | Acidic drugs                                  | Dialysis unit/SPE column                          | [13 |
|      |        | X    |        | X   | CE         | Blood serum              | Anti-IgE mAb                                  | Preconcentration-CE using an analyte concentrator | [24 |
|      |        | X    |        | X   | CE         | _                        | Methylamphetamine                             | Use of concentration step to collect analytes     | [25 |
|      |        | X    |        | X   | CE         | _                        | PAHs  | SPME with a optical fibre                         | [26 |
|      |        | X    |        | X   | CE         | Natural water            | Phenols                                       | SPME fibres                                       | [27 |
| K    |        |      |        | X   | CE         | Waste solids             | Alkylbenzenesulfonates                        | SPE with Carbopack B                              | [28 |
| (    |        |      |        | X   | CZE        | Fried meat and fish      | Heterocyclic aromatic amines                  | Clean up in cartridges                            | [29 |
|      |        |      | X      | X   | MEKC       | Chlorophenols            | Human urine                                   | MEKC coupled to a clean up system                 | [30 |
|      |        |      | X      | X   | CE         | Wine                     | Biogenic amines                               | C <sub>18</sub> adsorbent minicolumn              | [3] |
|      |        |      | X      | X   | CE         | Wines                    | trans-Resveratrol/polyphenols                 | C <sub>18</sub> SPE column                        | [32 |
|      |        |      | X      | X   | CE         | Inorganic cations        | Water   | Chelex-100 minicolumn incorporated in the CFS     | [33 |
|      |        |      | X      | X   | MEKC       | Pesticides               | Water   | C <sub>18</sub> SPE minicolumn was used           | [34 |
|      |        | X    |        | X   | CZE        | Pseudoephedrine          | Human plasma                                  | Combination of flow injection SPE and CZE         | [35 |
|      |        |      | X      | X   | CE         | Inorganic anions         | Water   | On-line ion-exchange in a flow injection-CE       | [36 |
|      |        | X    |        | X   | CE         | Serum and urine          | Sulfonamides                                  | Dialysis-SPE                                      | [67 |
|      |        | X    |        | X   | CE         | _                        | Collagen crosslinks HP and LP                 | SPE-dialysis LC system                            | [68 |
|      |        |      |        | X   | CGE        | Human plasma             | Phosphorothioate oligonucleotides             | SAX cartridge                                     | [80 |
|      |        |      |        |     | CE         | Environmental            | -   | Review  | [1] |
|      |        | X    |        | X   | CE         | Aqueous samples          | Environmental pollutants                      | Review  | [12 |
|      |        | Α.   |        | X   | CZE        | Plasma                   | E-5-(2 Brvinyl)-2'deoxyuridine                | SPE column  | [1: |
|      |        |      |        | X   | CE         | Urine                    | Catecholamines                                | SPE alumina cartridge                             | [12 |
|      |        | X    |        | X   | CE         | -                        | -   | Membrane preconcentration                         | [13 |
| (    |        | А    |        | X   | CE         | Water                    | Phenylurea herbicides                         | · · · · · · · · · · · · · · · · · · ·             | [12 |
|      |        |      |        | X   | CE         | Tap and river water      | Atrazine, terbutylazine                       | C <sub>18</sub> SPE<br>LiChrolut EN cartridges    | [13 |
|      |        |      |        | X   | MEKC       | Human urine              | Glucuronides of entacapone                    | Cartridges of Sep-Pak Vac C <sub>18</sub>         | [12 |
|      |        |      |        |     | CE         | Serum and urine          | Sulfonamides                                  | Cartridges packed                                 | -   |
|      |        | _    | X      | X   |            | Serum and urme           |   | • .   | [12 |
|      |        | X    |        | X   | CZE<br>CZE | Corr lumin and man must  | Fluorescein isothiocyanate<br>ein Polyphenols | SPE with polyomide contrides                      | [12 |
|      |        |      |        | X   |            | Soy, lupin and pea prote | **  | SPE with polyamide cartridges                     | [13 |
|      |        |      |        | X   | CE         | Soil                     | Imidazoline herbicides                        | C <sub>18</sub> cartridge                         | [1: |
|      | X      |      |        | X   | CE         | Proteins                 | Peptides                                      | Miniaturised reversed-phase C <sub>18</sub>       | [13 |
| (    |        |      |        | X   | CE         | Wines                    | cis- and trans-resveratrol                    | SPE cartridges                                    | [13 |
|      |        |      |        | X   | CE         | Seawater                 | Hydroxamate-type siderophores                 | SPE column  | [13 |
|      |        |      | X      | X   | CE         | Urine                    | Phenprocoumon (I)                             | SPE column  | [13 |
|      |        |      |        | X   | CE         | Human plasma and uri     |   | SPE column  | [1: |
|      |        |      |        | X   | CE         | Blood and urine          | Common illicit drug                           | GDX301 SPE  | [1: |
|      |        | X    |        | X   | CZE        | Bauhinia purpurea leci   |   | Adsorption preconcentration                       | [1  |
|      |        | X    |        | X   | CZE        | Tap water                | Haloacetic acids                              | Four types of adsorbents                          | [1: |
|      |        |      |        | X   | CE         | Rat urine                | Bupivacaine and its metabolites               | -   | [14 |
|      |        |      | X      | X   | CE         | Serum and urine          | Non steroid anti-inflammatory drugs           | LC-18 cartridges                                  | [14 |
|      |        |      |        | X   | CE         | Peptides substrate       | Chymotrypsin                                  | Streptavidin-agarose beads                        | [14 |
|      |        |      |        | X   | CE         | Potatoes and onions      | Maleic hydrazides                             | C <sub>18</sub> SPE cartridge                     | [14 |
|      |        |      |        | X   | CE         | Pea plant                | Indole-3-acetylaspartic acid                  | C <sub>18</sub> SPE cartridge                     | [14 |
|      |        |      |        | X   | CE         | Water                    | Aromatic sulfonates                           | LiChrolut EN SPE column                           | [14 |
| ι    |        |      |        | X   | CZE        | Plasma                   | Midazolam and its metabolites                 | Bond Elut C <sub>18</sub> cartridge               | [14 |
|      |        |      |        | X   | CZE        | Plasma                   | Methotrexate and leucovorin                   | Bakerbond C <sub>18</sub> SPE column              | [14 |
|      |        |      |        | X   | CGE        | Blood and plasma         | Antisense oligonucleotides                    | Cationic nanoparticles for SPE                    | [14 |
| (    |        |      |        |     | x CE       | Biological tissues-M     | Substances P                                  | Combination SPE-HPLC-CE                           | [14 |
| (    |        |      |        | X   | CE         | Haemophilus              | Pathogenic lipopolysaccharides                | C <sub>18</sub> column and membrane               | [15 |

Table 5. Continued

| Coup | ling n | node |        | CFS  | Electrophoretic | Sample                                   | Analytes                                | Remarks                                    | Ref.  |
|------|--------|------|--------|------|-----------------|--|---|--|-------|
| OFF  | IN     | ON   | OFF/ON | NC C | mode            |  |   |  |       |
| x    |        |      |        | х    | CE              | Drinking water                           | Pesticides                              | C <sub>18</sub> -bonded SPE cartridge      | [151] |
| X    |        |      |        | X    | CE              | Plasma                                   | Oligodeoxyribonucleotides               | -  | [152] |
| X    |        |      |        | X    | MEKC            | Milk                                     | Cyclopiazonic acid                      | Sep-Pak plus silica gel SPE                | [153] |
| X    |        |      |        | X    | CZE             | Foodstuffs                               | Histamine                               | Clean up of samples by SPE                 | [154] |
|      |        | X    |        | X    | CZE             | Proteins, proteomes                      | -                                       | SPE C <sub>18</sub> cartridge              | [155] |
|      |        | X    |        | X    | CE              | Insulin                                  | Somatomedin C, IGF I and IGF II         | Concentrator tips                          | [156] |
| X    |        |      |        | X    | CE              | Soybeans                                 | Multiple herbicides                     | SPE with Alumina-N dichloromethane         | [157] |
| X    |        |      |        | X    | MEKC            | Urine                                    | Nitrazepam, flunitrazepam, triazolam    | SPME                                       | [158] |
| X    |        |      |        | X    | CGE             | Human blood plasma                       | Phosphorothioate oligonucleotides       | Cationic polystyrene nanoparticles         | [159] |
| X    |        |      |        | X    | CE-MEKC         | Paddy water                              | Pesticides/Insecticides                 | SPE cartridges                             | [160] |
| X    |        |      |        | X    | CE              | Ground water                             | Sulfonated azo dyes                     | Cartridges of Isolute ENV                  | [161] |
| Κ    |        |      |        | X    | CE              | Groundwater                              | Fluorescein (I)                         | Styrene-divinylbenzene disks               | [162] |
| X    |        |      |        | X    | CZE-CEC         | Human urine                              | Metabolites of paracetamol              | C <sub>18</sub> SPE                        | [163] |
| X    |        |      |        | X    | CE              | Proteins                                 | Trypsin                                 | -  | [164] |
| X    |        |      |        | X    | CZE-ITP         | Proteins                                 | Tryptic digests of bovine serum albumin | Reversed-phase resins for SPE              | [165] |
| X    |        |      |        | X    | MEKC            | Water                                    | Sulfonated azo dyes                     | Cartridges Isolute ENV                     | [166] |
| X    |        |      |        | X    | CE              | Water                                    | Nitrophenols                            | C <sub>18</sub> membrane discs             | [167] |
| X    |        |      |        | X    | CE              | River water                              | Naphthalene sulfonates                  | Preconcentration by SPE                    | [168] |
| x    |        |      |        | x    | CE              | Human serum                              | Pentazocine enantiomers                 | Phenyl SPE cartridges                      | [169] |
| X    |        |      |        | X    | CZE             | Coffee cherries                          | Phloxine B and uranine                  | SPE column                                 | [170] |
| X    |        |      |        | X    | CZE             | Potable water                            | Hydroxymetabolites of atrazine          | Adsorbent cartridge                        | [171] |
| X    |        |      |        | X    | MEKC            | Ground water                             | Hexazinone and its metabolites          | Cartridge of Supelclean ENVI-Carb          | [172] |
| x    |        |      |        | X    | CE              | Laundry detergent                        | Alkylbenzenesulfonates                  | Clean up by SPE                            | [172] |
|      |        |      |        |      | CE              | Serum                                    |   | - ·  |       |
| X    |        |      |        | X    | CE              |  | R(+)-, $S(-)$ -pentobarbital            | C <sub>18</sub> SPE column                 | [174] |
| X    |        | _    |        | X    |                 | Human serum                              | Bile acids                              | Cartridges of Whatman ODS-3                | [175] |
|      |        | X    |        | X    | CE              | Sheep liver                              | Metallothioneins                        | Divinylbenzene resin SKP                   | [176] |
| X    |        |      |        | X    | MEKC            | Urine                                    | Corticosterone                          | SPME discs cartridges                      | [177] |
| X    |        |      |        | X    | CZE             | Juice                                    | Yellow and red safflower pigments       | ODS-4 cartridge                            | [178] |
| ζ.   |        |      |        | X    | CZE             | Human urine                              | β-Blocker atenolol                      | Bond Elut certify SPE columns              | [179] |
|      |        |      |        |      | CE              | Proteins                                 | -                                       | Review                                     | [180] |
|      |        |      |        |      | CE              | -  | Anions                                  | Membrane based SPE discs                   | [181] |
| X    |        |      |        | X    | CE              | Catfish                                  | Antibiotic oxytetracycline              | Sep-Pak C <sub>18</sub> SPE cartridges     | [182] |
| X    |        |      |        | X    | CE              | Rat serum                                | D-Pen 2,5 enkephalin                    | SPE cartridges                             | [183] |
| X    |        |      |        | X    | CE              | Plant tissue                             | Indole-3-acetic acid                    | SPE C <sub>18</sub> Bakerbond cartridges   | [184] |
| X    |        |      |        | X    | MEKC            | Human serum                              | Cortisone, dexamethasone                | SPE C <sub>18</sub> cartridges             | [185] |
| K    |        |      |        | X    | CE              | Human urine                              | Debrisoquine, 4-hydroxydebrisoquine     | C <sub>18</sub> cartridges                 | [186] |
| K    |        |      |        | X    | CE              | Urine                                    | Enantiomers 4-hydroxydebrisoquine       | Isolute ENV cartridges                     | [187] |
| X    |        |      |        | X    | CE              | Milk                                     | Tetracyclines antibiotics               | SPE MP1 micro-column                       | [188] |
| X    |        |      |        | X    | CE              | Human serum                              | Lipoproteins                            | C <sub>18</sub> Sep-Pak                    | [189] |
| X    |        |      |        | X    | CE              | Human plasma, urine                      | EDTA                                    | Anion-exchange disc                        | [190] |
| ζ    |        |      |        | X    | CE              | Water                                    | Sulfonylurea herbicides                 | Anion-exchange cartridges                  | [191] |
| ζ    |        |      |        | X    | CZE             | Blood serum                              | Proteins                                | Beckman Paragon SPE system                 | [192] |
| Κ    |        |      |        | X    | CZE             | Skin                                     | Fatty acid                              | Silica gel+LiChroprep                      | [193] |
| Κ    |        |      |        | X    | CE              | Oral solid dosage                        | Betamethasone (I), ergotamine tartrate  | Empore C <sub>18</sub> SPE discs           | [194] |
| X    |        |      |        | X    | CE              | River water                              | Naphthalenesulfonates                   | _  | [195] |
| x    |        |      |        | X    | CZE             | Plasma                                   | Linear pentapeptide dolastin            | Sep-Pak C <sub>18</sub> SPE column         | [196] |
| X    |        |      |        | X    | CZE             | Clinical urine                           | Orotic acid                             | RP-18 SPE column                           | [197] |
|      |        |      |        | x    | CZE             | Beet, tobacco, wheat                     | Cytokinins                              | SPE on C <sub>18</sub> cartridges          | [198] |
| X    |        |      |        | X    | CE              | Human serum                              | S(+)-, $R(-)$ -ondansetron              | SPE on cyanopropyl cartridge               | [199] |
|      |        |      |        | X    | CE              | Urine and serum                          | Barbiturates                            | SPME device                                | [200] |
| X    |        |      |        | Λ    | CE              | Tryptic digest from bovine serum albumin |   | Fused silica column C <sub>18</sub> silica | [200] |

Table 5. Continued

| Coupl  | ling m | ode |        | CFS    | Electrophoretic | Sample                       | Analytes                                 | Remarks  | Ref.  |
|--------|--------|-----|--------|--------|-----------------|------------------------------|--|--|-------|
| OFF    | IN     | ON  | OFF/ON | NC     | C mode          |                              |  |  |       |
| x      |        |     |        | х      | CE              | Shellfish tissues            | Domoic acid                              | LC-SAX cartridge                               | [202] |
| X      |        |     |        | X      | CE              | Plant tissue                 | Indol-3-yl acetic acid                   | C <sub>18</sub> SPE column                     | [203] |
| X      |        |     |        | X      | CE              | Water                        | Aliphatic amines                         | Cartridges and discs                           | [204] |
| X      |        |     |        | X      | CE              | Environmental water          | Ethylenediaminetetraacetic acid          | Anion-exchange disc                            | [205] |
| X      |        |     |        | X      | CE              | Human or rat urine           | Paracetamol and phenacetin               | IST C <sub>18</sub> bonded cartridges          | [206] |
| X      |        |     |        | X      | CZE             | Human urine                  | Enantiomers of methadone (I)             | SPE on Bond Elut cartridge                     | [207] |
| X      |        |     |        | X      | CE              | Water                        | Phenolic compounds                       | ENVI-Chrom P cartridge                         | [208] |
| X      |        |     |        | X      | CZE             | Neonatal urine               | Lactate, pyruvate, organic acids         | C <sub>18</sub> SPE cartridges                 | [209] |
| X      |        |     |        | X      | CZE-MEKC        | Blackcurrant bud extract     | Flavonoids, cinnamic, phenolic acid      | Celite SPE column                              | [210] |
| X      |        |     |        | X      | MEKC            | Human plasma                 | Fluconazole                              | Bakerbond C <sub>18</sub>                      | [211] |
| X      |        |     |        | X      | MEKC            | Equine urine                 | Morphine and meclofenamic acid           | Bond Elut C2 column                            | [212] |
|        | X      |     |        | X      | AGE             | Human serum                  | Proteins                                 | Prepoured REP-agarose gel                      | [213] |
| X      |        |     |        | X      | CE              | Biological matrices          | Drug identification                      | Accubond Evidex cartridge                      | [214] |
| X      |        |     |        | X      | MEKC            | Jelly and juice              | Anthraquinone pigments                   | _  | [215] |
| X      |        |     |        | X      | CZE             | Plant tissues                | Ascorbate                                | C <sub>18</sub> SPE cartridge                  | [216] |
|        |        | X   |        | X      | CE              | _                            | Metal ions                               | SPE cartridge                                  | [217] |
| X      |        |     |        | X      | CZE             | Tobacco                      | Carbaryl and carbofuran                  | Florisil SPE cartridge                         | [218] |
| X      |        |     |        | X      | CZE             | Rats                         | [p-penicillamine 2,5]enkephalin in       | C <sub>18</sub> SPE column                     | [219] |
| X      |        |     |        | X      | CZE             | _                            | Eleven priority phenols                  | Styrene-divinylbenzene                         | [220] |
| X      |        |     |        | X      | CE              | Human serum                  | Benzodiazepines                          | Cleaned-up by SPE                              | [221] |
| X      |        |     |        | X      | CZE-ITP         | Urine                        | Adenosine                                | Sep-Pak SPE column                             | [222] |
| X      |        |     |        | X      | CE              | Water                        | Trialkylstannane derivates               | XAD-2 column                                   | [223] |
|        |        | X   |        | X      | CE              | _                            | Peptides                                 | C <sub>18</sub> coupled to silica capillary    | [224] |
|        |        | X   |        | X      | CZE             | Drugs                        | Hypoglycaemic sulfonylurea               | Bakerbond C <sub>18</sub> SPE silica           | [225] |
| X      |        |     |        | X      | CE              | Rat liver                    | Lycopus europaeus L.                     | Bakerbond Phenyl column                        | [226] |
| X      |        |     |        | X      | CE              | Urine                        | Enantiomers of amphetamines              | Bond Elut certify cartridge                    | [227] |
| X      |        |     |        | X      | CZE             | Rain water                   | Heterocyclic aromatic amines             | Bond Elut C <sub>18</sub> SPE cartridge        | [228] |
| X      |        |     |        | X      | MEKC            | Human urine                  | Hypoglycaemic drugs                      | Samples extracted by SPE                       | [229] |
|        |        |     |        |        | CE              | Biological fluids            | Neuropeptides                            | Review   | [230] |
| X      |        |     |        | X      | CZE             | Water                        | Aromatic amines                          | Cartridge of copolymer                         | [231] |
| X      |        |     |        | X      | CE              | Urine                        | Ibuprofen, fluriprofen, aspirin          | C <sub>18</sub> SPE cartridge                  | [232] |
| X      |        |     |        | X      | CE              | Paper                        | Anion                                    | SPE as a sample clean up                       | [233] |
| X      |        |     |        | X      | CE              | Soil                         | Sulfonylurea herbicides                  | C <sub>18</sub> SPE columns                    | [234] |
| X      |        |     |        | X      | CZE             | Human plasma                 | Cimetidine                               | Supelclean LC-18 SPE                           | [235] |
| X      |        |     |        | X      | CZE             | Household detergents         | Alkylbenzenesulfonates                   | ODS C <sub>18</sub> SPE column                 | [236] |
| X      |        |     |        | X      | CE              | Rat liver microsomes         | Theophylline and their metabolites       | SPE on C <sub>18</sub> columns                 | [237] |
|        |        | X   |        | X      | CE              | Bovine serum albumin digests | Peptides                                 | Reversed-phase C <sub>18</sub> packing         | [238] |
| X      |        |     |        | X      | CE-ITP          | Plasma                       | Heterocyclic peptides                    | SPE cartridge                                  | [239] |
| X      |        |     |        | X      | CE              | Biological fluids            | Organic anions                           | _  | [240] |
| X      |        |     |        | X      | CZE             | Urine                        | Organic acids                            | C <sub>18</sub> SPE cartridge                  | [241] |
| X      |        |     |        | X      | CE              | Wines                        | Organic acids                            | C <sub>18</sub> SPE cartridge                  | [242] |
| X      |        |     |        | X      | CE              | DNA adduct                   | Nucleotides                              | Analytes separated by SPE                      | [243] |
| X      |        |     |        | X      | CZE             | Urine                        | Racemethorphan and racemorphan           | BakerBond Octadecyl HC                         | [244] |
| X      |        |     |        | X      | CZE             | Water                        | Phenoxy acid herbicides                  | C <sub>18</sub> membrane extraction (Overview) | [244] |
| X      |        |     |        | X      | MEKC            | Urine                        | 9-Tetrahydrocannabinol-9-carboxylic acid | -  | [245] |
|        |        |     |        | X      | ITP             | Blood serum                  | γ-Aminobutyric acid                      | Separcol Si-C <sub>18</sub> L minicolumn       | [247] |
| X<br>v |        |     |        |        | MEKC            | Serum                        | Cimetidime                               | Reversed-phase C <sub>18</sub>                 | [248] |
| X<br>v |        |     |        | X<br>v | CE              | Human serum                  | Cytarabine                               | C <sub>18</sub> SPE cartridges                 | [246] |
| X      |        |     |        | X      | MEKC            | Tablets                      | Water soluble vitamins                   | Octadecylsilane SPE column                     |       |
| X      |        |     |        | X      |                 |                              |  |  | [385] |
| X      |        |     |        | X      | CE              | Lung tissue                  | Angiotensins                             | C <sub>18</sub> extraction cartridge           | [387] |

CFS: Continuous flow system; NC: no chromatography; C: chromatography; CZE: capillary zone electrophoresis; CE: capillary electrophoresis; MEKC: micellar electrokinetic chromatography; SPE: solid-phase extraction; SPME: solid-phase microextraction; CGE: capillary gel electrophoresis; CEC: capillary electrochromatography; ITP: isotachophoresis; AGE: agarose gel electrophoresis.

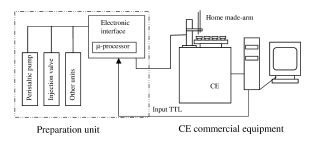


Fig. 2. Continuous flow system-capillary electrophoresis, commercial equipment (adapted from Ref. [34]).

using direct UV absorbance detection was developed to demonstrate the usefulness of this arrangement (see Fig. 3) [36].

### 9. Preconcentration membrane systems coupled to capillary electrophoresis

Poor concentration limits of detection (LODs) of CE methods often preclude their use for the analysis of dilute analyte mixtures. This limitation was addressed by the development of analyte concentrator and membrane preconcentration cartridges. The development of membrane preconcentration was undertaken to decrease or remove all the limitations observed in studied of solid-phase preconcentration [37,38]. The membranes have been used in more than 90 references shown in Table 6.

Using a suitably coated/impregnated membrane it is possible to minimise the bed volume of adsorptive phase at the inlet of the preconcentration capillary. The membrane is installed in a cartridge that is

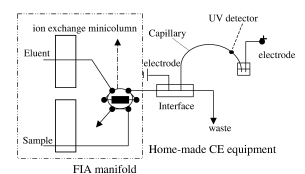


Fig. 3. Flow system-capillary electrophoresis, laboratory-made (adapted from Ref. [36]).

usually prepared from PTFE tubing. In addition to analyte preconcentration, membranes can also be used to effect sample clean up. This is particularly important for physiologically derived samples such as blood, urine, etc., where the presence of high salt concentrations can dramatically effect analyte separation by CE.

Small lengths of narrow-bore tubular membrane [39] can be interposed in the separation capillary in CE separation systems. These membrane segments can be used as sampling interfaces; a jacket is built outside the membrane, and the sample is introduced by diffusion/permeation through the membrane. Various examples are shown; the determination of gaseous samples through a porous membrane, the determination of ionisable/nonionic solutes by permeation through a silicone rubber membrane, and the separation of low-molecular-mass constituents in blood plasma by transport through a dialysis membrane.

In 1996, Szostek and Koropchak [40] described two means for interfacing condensation nucleation light scattering detection to CE. With the first method, a fused-silica capillary was used for the separation and the CE was grounded through a Nafion membrane that also connected the system to a microconcentric pneumatic nebuliser.

In the same year, a CE post-column radionuclide detector was developed that uses a commercial phosphor-imaging detector and was optimised for low-energy  $\beta$  emitters. Eluent from the separation capillary was deposited on a membrane. Emission from radioactive analytes on the membrane was integrated using the phosphor-imaging detector system. Results from the phosphor-imaging system were converted to conventional electropherograms [41].

A hollow fibre miniaturised supported liquid membrane (SLM) device for sample preparation is connected on-line with CE and used for determination of a basic drug, bambuterol, in human plasma. The analyte was extracted from the outside of the hollow fibre (donor) through the liquid membrane (pores of the fibre impregnated with organic solvent) into the acceptor solution in the fibre lumen. Very clean extracts of low ionic strength are obtained from the SLM treatment, making this sample pretreatment method compatible with the CZE double-stacking procedure, which in turn makes it possible to inject

Table 6 Coupling continuous membrane systems to capillary electrophoresis

| Coup | ling mo | ile      | CFS  | Electrophoretic | Sample                             | Analytes                                     | Remarks  | Ref. |
|------|---------|----------|------|-----------------|------------------------------------|--|--|------|
| OFF  | IN O    | N OFF/ON | NC C | mode            |                                    |  |  |      |
|      | Х       |          | X    | CE              | -                                  | Proteins                                     | Interface based on microdialysis memebrane         | [10] |
| X    |         |          | X    | CE              | -                                  | Peptides                                     | Preconcentration cartridge contained a membrane    | [37] |
|      | X       |          | X    | CE              |                                    | Peptides                                     | Membrane preconcentration                          | [38] |
|      | X       |          | X    | CZE             | Blood plasma                       | Low-molecular-mass analytes                  | Membranes were used as sampling interfaces         | [39] |
|      | X       |          | X    | CZE             | _                                  | Acids and peptides                           | Capillary grounded electrically through a membrane | [40] |
|      | x       |          | x    | CE              | -                                  | Radionuclides                                | Eluate was deposited on to a membrane              | [41] |
|      | X       |          | X    | CZE             | Human plasma                       | Basic drugs                                  | Supported liquid membrane                          | [42] |
|      | x       |          | x    | CE              | Beverage                           | Carbonate                                    | Gaseous products trough a membrane                 | [43] |
| ζ    |         |          | x    | CZE             | _                                  | Amino acids                                  | Concentration using liquid membranes               | [44] |
|      |         |          |      | CE              | Blood plasma                       | Total and unbound drug                       | Review   | [63] |
|      | х       |          | x    | IEF             | Pharmalyte 5-8                     | Carbonic anhydrase                           | Separated by cellulose membrane                    | [70] |
| ζ    |         |          | X    | CE              | Serum                              | Oligonucleotides                             | Membrane filters                                   | [71] |
| ζ.   |         |          | X    | CE              | Blood plasma                       | Oligonucleotides                             | Dialysis using a cellulose membrane                | [72] |
| ζ.   |         |          | X    | CE              | Polymerase chain reaction products | Nucleic acids                                | Membrane dialysis was a sample clean up            | [77] |
|      |         |          | X    | CE              | Smoked fish                        | Clostridium botulinum                        | Membrane dialysis before analysis                  | [78] |
| ζ.   |         |          | X    | CE              | DNA fragments                      | Clostridium botulinum                        | Products were desalted by membrane dialysis        | [79] |
| ζ.   |         |          | X    | CGE             | Plasma                             | Phosphorothioate                             | Dialysis with a Millipore VS membrane              | [80] |
|      |         |          | X    | CE              | Living rat brain                   | 4-Aminobutyric acid                          | Polycarbonate membrane dialysis probe              | [81] |
| ζ.   | v       |          |      | CE              | Living fat brain                   | 4-Ammobutyric acid                           | Membrane preconcentration                          | [124 |
| ,    | X       |          | X    | CE              | Seawater                           | Cidorophoros                                 | Membrane filter                                    | [134 |
|      |         |          | X    |                 |                                    | Siderophores  Pothogonia limonalysasaharidas |  | -    |
|      | X       |          | Х    | CE              | Haemophilus                        | Pathogenic lipopolysaccharides               | On-line preconcentration                           | [150 |
|      |         |          |      | CE              | Proteins                           | -  | Review   | [180 |
|      |         |          | Х    | CE              | -                                  | Anions                                       | SPE discs comprise PTFE membrane                   | [18] |
|      |         |          | X    | CE              | Paper                              | Anion  | Membrane based on SPE as a sample clean up         | [233 |
| (    |         |          | X    | ITP             | -                                  | Gaseous sulfur dioxide                       | Membrane collector                                 | [250 |
| (    |         |          | X    | CE              | -                                  | Trace metals                                 | Hollow fibre membrane                              | [25] |
|      | X       |          | X    | CZE             | -                                  | Inorganic anions                             | Membrane separated                                 | [25] |
|      | X       |          | X    | CIEF            | -                                  | -  | Membrane sample preparation                        | [253 |
| K    |         |          | X    | CE              | Rat                                | Monoamine and their metabolites              | Polycarbonate ether membrane                       | [254 |
| K    |         |          | X    | IEF-CE          | Pharmaceutical                     | Erythropoietin glycoforms                    | Centrifuging through a membrane                    | [255 |
|      | X       |          | X    | CE-FI           | Drug dissolution                   | Trimethoprim, sulfamethoxazole               | On-line membrane filter                            | [256 |
| (    |         |          | X    | CZE             | Blood plasma                       | Human IgG                                    | Affinity membrane cartridge                        | [257 |
| (    |         |          | X    | CGE             | Plants                             | RNA  | Positively charged nylon membrane                  | [258 |
| ζ    |         |          | X    | MEKC            | Pharmaceutical                     | Amphotericin B                               | Filtration through membrane filters                | [259 |
| Κ    |         |          | X    | CE              | -                                  | Organic ions, alkylamines                    | Poly(vinyl chloride) membranes                     | [260 |
| (    |         |          | X    | CE              | Cellulose solutions                | Supercoiled plasmid DNAs                     | Float membrane                                     | [26] |
|      |         |          | X    | CE              | -                                  | Proteins                                     | Membrane preconcentration                          | [26  |
|      |         |          | X    | CZE             | Feed solutions                     | Humic acids                                  | Polyethersulfone cut-off membrane                  | [26  |
| (    |         |          | X    | CE              | Medicines                          | Magnolol and honokiol                        | Filter membrane                                    | [26  |
|      |         |          | X    | CE              | _                                  | DNA sequence                                 | Ultrafiltration trough a polymembrane              | [26: |
|      | X       |          | x    | CE              | _                                  | Proteins                                     | Membrane preconcentration                          | [260 |
|      |         |          | X    | CE              | _                                  | Dansylated amino acids                       | Membrane collection                                | [26] |
|      |         |          | X    | CE              | Tumour peptide                     | Particular antigen                           | Membrane mounted in a length of silica tubing      | [268 |
|      |         |          | X    | CE              | -                                  | Peptides                                     | Membrane preconcentration                          | [269 |
|      |         |          | X    | CE              | Cod muscle tissue                  | $M_{\rm r}$ 41 000 protein                   | Membrane filter                                    | [270 |
|      |         |          | X    | CE              | _                                  | Amino acids                                  | Liquid membrane extraction                         | [27] |
|      |         |          | X    | CIE             | Pharmaceutical                     | Inorganic and organic anions                 | PVDF membrane filters                              | [272 |
|      |         |          | X    | CE              | Pericarpium                        | Alkaloids                                    | Filter membrane                                    | [273 |
| (    | v       |          | X    | CZE             |                                    | Phenylacetate inositol                       | Separated by cellulose membrane                    | [274 |
|      | X       |          | Λ    | CLL             | _                                  | i nenyiacetate mositor                       | separated by certaiose memorane                    | [4/4 |

Table 6. Continued

| Coupl | ing mo | ode |        | CFS |   | Electrophoretic | Sample               | Analytes                         | Remarks   | Ref  |
|-------|--------|-----|--------|-----|---|-----------------|----------------------|----------------------------------|---|------|
| OFF   | IN     | ON  | OFF/ON | NC  | С | mode            |                      |                                  |   |      |
| x     |        |     |        | х   |   | CE              | Water                | Selenium and arsenic             | Gaseous hydrides trough membrane                            | [27  |
| K     |        |     | X      |     |   | CZE             | Minced fish          | Histamine                        | Filtered through a membrane                                 | [27] |
| K     |        |     | X      |     |   | CZE             | Blood plasma         | Inositol phosphates              | Separated by two membranes                                  | [27  |
| ζ     |        |     | X      |     |   | CE              | _                    | Substance P and its metabolism   | Membrane reactor  | [27  |
| (     |        |     |        | X   |   | CGE             | Plasma               | Proteins                         | Nylon membrane  | [28  |
| (     |        |     |        | X   |   | CZE             | Vegetables           | Ascorbic acids                   | Filtered through a membrane filter                          | [28  |
| (     |        |     |        | X   |   | CZE             | Blood serum          | Proteins                         | Comparison between CZE and CAME                             | [28  |
|       |        |     |        | X   |   | CZE             | Synthetic food       | 2-Nitrophenol                    | Movement of the membrane                                    | [28  |
| (     |        |     |        | X   |   | CE              | Drinking water       | Arsenic compounds                | Membrane gas-liquid separator                               | [28  |
|       |        |     |        | X   |   | CE              | Peat                 | Fulvic acids                     | Dialyzed with a cellulose ester membrane                    | [28  |
| (     |        |     |        | X   |   | CE              | Blood plasma         | Paracetamol, salicylic acid      | Plasma was filtered through a membrane filter               | [28  |
|       |        |     | X      |     | X | CZE             | Plasma               | Bambuterol                       | Sample enriched by passing through a SLM                    | [28  |
| (     |        |     |        | X   |   | CE              | Labile cyanide       | Metallocyanides                  | Concentration on supported liquid membranes                 | [28  |
|       |        | X   |        | X   |   | CZE             | Liquors              | Adenosine triphosphate           | Electrodialysis system separated by a membrane              | [28  |
|       |        | X   |        | X   |   | CE              | Biological materials | Peptides and proteins            | Membrane preconcentration                                   | [29  |
|       |        |     |        | X   |   | MEKC            | Human blood          | Creatinine                       | Centrifuged through a membrane                              | [29  |
|       |        |     |        | X   |   | CZE             | Vegetables           | Free calcium                     | Filtered through a membrane                                 | [29  |
| (     |        |     |        | X   |   | CE              | Blood plasma         | Bambuterol                       | Extracted through a porous PTFE membrane                    | [29  |
|       | X      |     |        | X   |   | ITP             | _                    | Peptides                         | Preconcetration cartridge incorporating a membrane          | [29  |
|       | X      |     |        | X   |   | CE              | _                    | Organic acids                    | Liquid membrane phases                                      | [29  |
| (     |        |     |        | X   |   | CZE             | Human milk           | Organometallic compounds         | Milk was ultrafiltrated through a membrane                  | [29  |
|       |        |     |        | X   |   | CGE             | _                    | Proteins                         | Miniature ultrafiltration sampling probe (membrane)         | [29  |
| •     |        | X   |        | X   |   | CE              | EL-4 cells           | Peptides                         | Membrane preconcentration                                   | [29  |
|       | х      |     |        | x   |   | CE              | Biological material  | Biological compounds             | Membrane installed in the cartridge of a capillary          | [29  |
|       | Α      |     |        | X   |   | CE              | Food                 | Histamine                        | Membrane filtration analysis before analysis                | [30  |
|       |        |     |        | X   |   | CE              | Hepatocytes          | 3-Phenylaminopropane             | Membrane was used for sample concentration                  | [30  |
|       |        |     |        | X   |   | MEKC            | Cheddar cheese       | Caseins and large peptides       | Buffer fractionated through a cutoff membrane               | [30  |
|       |        |     |        | X   |   | CZE             | Plasma               | Basic drugs                      | Supported liquid membrane technique                         | [30  |
|       |        | X   |        | X   |   | CE              | -                    | Haloperidol/peptides             | Membrane preconcentration                                   | [30  |
|       |        | X   |        | X   |   | CE              | _                    | Peptides                         | Preconcentration membrane                                   | [30  |
|       |        | А   |        | A   |   | CE              | _                    | Peptides Peptides                | Solid-phase reactor for preconcentration                    | [30  |
| ,     |        |     |        | X   |   | CE              | Biological fluids    | Phosphorothioate oligonucleotide | Portions of the HPLC fractions were dialysed                | [30  |
| ,     |        |     |        | X   |   | CE              | Red blood cells      | Glutathione                      | Solution was filtered through a membrane                    | [30  |
|       |        |     |        |     |   | CE              | –                    |                                  | S .   | [30  |
|       |        | X   |        | X   |   | CE              |                      | Peptides and amino acids         | Eluent was directed on to peptide-binding membrane          | -    |
|       |        |     |        | X   |   |                 | Rat retina           | Enkephalin                       | The samples were centrifuged through a membrane             | [3]  |
|       |        |     |        | X   |   | CE              |                      | Proteins                         | Blotting membrane micro-preparation                         | [3]  |
|       | X      |     |        | X   |   | CZE             | Wheat flour          | Calcium and magnesium            | Liquid membrane (borax, ethylene glycol, EDTA)              | [3]  |
|       |        |     |        |     |   | CZE             | Environmental        | Phenoxy acid herbicides          | Use of C <sub>18</sub> membrane extraction discs (overview) | [24  |
|       | X      |     |        | X   |   | CE              | -<br>-               | Cations and anions               | Suppressors were of ion-exchange membrane                   | [31  |
|       |        |     |        | X   |   | CZE             | Foodstuffs           | Additives                        | Filtered through a micromembrane                            | [31  |
|       |        | X   |        | X   |   | CE              | Food                 | Anions                           | Capillary is linked to a cation-exchange membrane           | [3]  |
|       |        | X   |        | X   |   | CE              | -                    | Proteins                         | Membrane assembly at the exit of a capillary                | [31  |
|       | X      |     |        | X   |   | CE              | Blood serum          | Proteins                         | Fractions were collected on a moving membrane               | [31  |
|       | X      |     |        | X   |   | ITP             | Flowers              | Flavonoids                       | The filtrate was injected through the inlet membrane        | [31  |
| X     |        |     |        | X   |   | CZE             | Serum                | Lithium                          | Samples were deproteinised with a filter membrane           | [31  |

CFS: Continuous flow system; NC: no chromatography; C: chromatography; CZE: capillary zone electrophoresis; CE: capillary electrophoresis; SPE: solid-phase extraction; MEKC: micellar electrokinetic chromatography; CGE: capillary gel electrophoresis; ITP: isotachophoresis; IEF: capillary isoelectric focusing; CIE: capillary ion electrophoresis; SLM: supported liquid membrane.

large volumes of sample onto the separation capillary [42].

On-line gas diffusion was coupled to a laboratory-made CE system via a specially designed interface [43]. The sample was merged with a modifying solution, e.g., a strong acid, in a flow system to transform the analytes of interest into their respective gaseous forms. These transformed, gaseous analytes permeate through a PTFE membrane into an acceptor stream comprising of a Tris buffer.

Finally, among others, basic studies of a procedure for extraction of amino acids using a supported liquid membrane were presented by Wieczorek et al. [44]. The extractions were made from an aqueous donor phase with pH 3 to a more acidic acceptor phase and the mass transfer was driven by the proton gradient between these phases. Both acceptor solutions were analysed by CE.

### 10. Extraction coupled to capillary electrophoresis

Medina et al. [45] developed a method for speciation of organomercurials in marine samples using CE. Organomercurials were extracted from the samples by the Westoo procedure. The cysteine-organomercury complexes formed were separated by CE. The mercury species, ethylmercury, methylmercury, phenylmercury and inorganic mercury were well resolved in 12 min. The procedure was applied to the analysis of dogfish muscle certified reference material, mussel, cockle, clam and tuna and the results were compared with those obtained by GC.

Liu and Sheu [46] carried out the determination of the six major flavonoids in Scutellariae Radix by MEKC. Scutellariae Radix (SR) is the root of *Scutellaria baicalensis* Georgi and is commonly used in Chinese herbal drugs. Pulverised SR (0.1 g) was extracted with 7.5 ml of aq. 50% ethanol by reflux for 30 min and centrifuged (1500 g) for 5 min. Extraction was repeated three times. The solution was analysed on a fused-silica capillary.

Humic acid was separated into two fractions (A and D) by CZE by Rigol et al. [47]. Humic acids are normally extracted from the soil using diluted aqueous alkaline solutions. Sodium hydroxide has been widely used, but there are undesirable features such

as autoxidation or condensation of organic constituents. These changes can be minimised by performing extraction under an  $N_2$  atmosphere (see Table 7).

### 11. Isotachophoresis coupled to capillary electrophoresis

Electrophoresis-based techniques such as isotachophoresis (ITP) can also be used for sample preparation prior to CE analysis (see Table 8).

ITP as an on-line concentration pretreatment technique in CE was described by Stegehuis et al. [48]. On-line coupling of ITP with CE was studied as a means of lowering the range of application for the latter technique. Detection limits could be improved by ≥2 orders of magnitude and the additional selectivity of the combined system gave promising results concerning biological samples. For proteins, the electromigration characteristics are hardly changed by isotachophoretic pretreatment. The potential of the combined system was illustrated by separation ion of phthalaldehyde and fluorescein isothiocyanate derivatives of amino acids. A limitation of the coupled system is that a compromise concerning the buffer is needed between analysis time and resolution in each part of the system.

Foret et al. [49] carried out a trace analysis of proteins by CZE with on-column transient isotachophoretic preconcentration. The sample was concentrated by ITP using 0.01 M acetic acid as terminating anode electrolyte for 4 min. Sharp and well-resolved peaks were detected due to the initial ITP focusing. On-column transient and coupled-column isotachophoretic preconcentration of protein samples in CZE was developed by the same authors [50]. By this method, the conventional single-column instrument could be used for isotachophoretic sample preconcentration of 50-fold or more without any modification. The same protein mixture was used to demonstrate a coupled-column system, which provides more freedom in selecting capillary zone electrophoretic running conditions, the possibility of injecting higher sample volume, effective sample clean up and selective ion analysis.

Analyte focusing in CE using on-line ITP was carried out by Stegehuis et al. [51]. Derivatised

Table 7 Coupling continuous extraction systems to capillary electrophoresis

| Coupl  | ing mo | ode |        | CFS |   | Electrophoretic | Sample                | Analytes                         | Remarks  | Ref.  |
|--------|--------|-----|--------|-----|---|-----------------|-----------------------|----------------------------------|--|-------|
| OFF    | IN     | ON  | OFF/ON | NC  | С | mode            |                       |                                  |  |       |
| x      |        |     |        | Х   |   | CZE             | Lichens               | Cations                          | Lichen immersed in an extraction tube  | [4]   |
|        |        |     | X      | X   |   | CE              | Tea                   | Polyphenols                      | Sample preparation: extraction+filtration  | [5]   |
| X      |        |     |        | X   |   | MEKC            | Swabs                 | Organic gunshot                  | Extraction was by ultrasonic agitation   | [17]  |
| X      |        |     |        | X   |   | CZE             | _                     | Amino acids                      | Extraction of the amino acids from the donor   | [44]  |
| X      |        |     |        | X   |   | CE              | Marine samples        | Speciation of organomercury      | Extraction by the Westoo procedure   | [45]  |
| X      |        |     |        | X   |   | MEKC            | Chinese medicines     | Flavonoids                       | Extraction with ethanol  | [46]  |
| X      |        |     |        |     |   | CZE             | Soil                  | Humic acid                       | Extraction with NaOH and Na <sub>4</sub> P <sub>2</sub> O <sub>7</sub>                                 | [47]  |
| X      |        |     |        | X   |   | CE              | Urine                 | Tramadol                         | SPE  | [97]  |
| X      |        |     |        | X   |   | MEKC            | Seed meals of citrus  | Limonoid glucosides              | Soxhlet extracted to remove oils   | [105] |
|        |        |     |        |     |   | CE              | Environmental samples | _                                | Review (techniques for pretreatment)   | [117] |
| X      |        |     |        | X   |   | CE              | Antioxidative         | Plant beverages                  | Soxhlet extraction   | [120] |
| x      |        |     |        | x   |   | CE              | Pea plant             | Indole-3-acetylaspartic acid     | Analytes were extracted during 1 h.  | [144] |
| x      |        |     |        | X   |   | CE              | Body fluids           | Midazolam and its metabolites    | Liquid-liquid extraction and SPE   | [146] |
| X      |        |     |        | x   |   | CE              | Foodstuffs            | Histamine                        | Clean up by liquid–liquid extraction   | [154] |
| X      |        |     |        | X   |   | CE              | Soya bean             | Herbicides                       | Pressurised liquid extraction in acetonitrile  | [157] |
| X      |        |     |        | X   |   | CZE             | Beet, tobacco, wheat  | Cytokinins ribosides             | Extraction with <i>n</i> -butanol and SPE  | [198] |
| X      |        |     |        | X   |   | CE              | Water                 | Aliphatic amines                 | Ion pairing with extraction discs  | [204] |
|        |        |     |        | X   |   | CE              | Water                 | EDTA                             |  | [204] |
| X<br>X |        |     |        | X   |   | MEKC            | Human plasma          | Fluconazole                      | Strong anion-exchange extraction disc<br>Liquid-liquid extraction with CH <sub>2</sub> Cl <sub>2</sub> | [211] |
|        |        |     |        |     |   | CE              | Human serum           | Benzodiazepines                  |  | [221] |
| X      |        |     |        | X   |   | CE              |                       | •                                | Clean up by solvent extraction and SPE   |       |
| _      |        |     |        | _   |   |                 | Biological fluids     | Neuropeptides                    | Review (sample preparation method)   | [230] |
| X      |        |     |        | X   |   | CZE             | Water/soil            | Aromatic amines                  | SPE-liquid-liquid extraction   | [231] |
| X      |        |     |        | X   |   | CE              | Biological fluids     | Organic anions                   | Six zwitterionic free acids used for extraction  | [240] |
| X      |        |     |        | X   |   | CE              | Sclerotia             | Ergot alkaloids                  | Liquid extraction (twice)  | [246] |
| X      |        |     |        | X   |   | CE              | Po.                   | Amino acids                      | Liquid membrane extraction   | [271] |
| X      |        |     |        | X   |   | CE              | Plasma                | Bambuterol                       | Supported liquid membrane for pretreatment   | [293] |
| X      |        |     |        | X   |   | CZE             | Plasma                | Basic drugs (bambuterol)         | Extraction in supported liquid membrane  | [303] |
| X      |        |     |        | X   |   | CE              | Plasma/urine          | E-5-2-Bromovinyl-2'-deoxyuridine | Extraction buffer for determination of analyte   | [320] |
|        |        |     |        |     |   | CE              | Drug                  | _                                | Overview (FI solvent extraction system)  | [321] |
| X      |        |     |        | X   |   | CE              | Wheat flour protein   | Puroindolines                    | Extraction of soluble proteins   | [322] |
| X      |        |     |        | X   |   | MEKC            | Food                  | Biogenic amines                  | Amines were extracted with an acid solution  | [323] |
|        |        |     |        |     |   | CE              | Food/beverages        | Inorganic ions                   | Review (discussion of extraction method)   | [324] |
| X      |        |     |        | X   |   | CZE             | Roast coffee          | Hydroxycinnamic acids            | Extraction with organic solvent  | [325] |
| X      |        |     |        | X   |   | CZE             | Lettuce               | Vitamin C and inorganic cations  | Extraction with water and oxalic acid  | [326] |
| X      |        |     |        | X   |   | CGE             | Blood                 | Apolipoprotein E genotypes       | DNA was extracted by two different methods   | [327] |
|        |        |     |        |     |   | CE              | Food                  | Contamination analytes           | Review (examples of extraction techniques)   | [328] |
| X      |        |     |        | X   |   | CE              | Medicines             | Aconitine alkaloids              | Extraction with different solvents   | [329] |
| X      |        |     |        | X   |   | CE              | Foetal sheep liver    | Metallothionein isoforms         | Samples subjected to a two step extraction   | [330] |
| X      |        |     |        | X   |   | CE              | Drugs                 | Naphazoline, dexamethasone       | Sample was analysed after acetone extraction   | [331] |
| X      |        |     |        | X   |   | CZE             | Human erythrocytes    | Enzyme activities                | Diethyl ether extraction   | [332] |
| X      |        |     |        | X   |   | MEKC            | Paper food            | Biocides                         | Extraction with hot water  | [333] |
|        |        | X   |        | X   |   | CE-CGE          | DNA                   | HIV-I diagnosis                  | DNA extraction from samples on line  | [334] |
| X      |        |     |        | X   |   | CE              | Scleratia             | Ergot alkaloids                  | Extraction with a mixture of solvent   | [335] |
| x      |        |     |        | X   |   | CZE             | Water                 | Haloacetic acids                 | Preconcentration by liquid-liquid extraction   | [336] |
|        |        |     |        |     |   | CE              | _                     | Metallothionein isoforms         | Review   | [337] |
| X      |        |     |        | X   |   | CE              | Water                 | Phenols                          | Adsorption of phenols on organo-clay   | [338] |
|        |        |     | X      | X   |   | CE              | Soil                  | Anions                           | Automated extraction/filtering   | [339] |
| x      |        |     |        | X   |   | CE              | Puerariae radix       | Herbal medicinal components      | Threefold methanol extraction  | [340] |
| X      |        |     |        | X   |   | MEKC            | Liver and kidney      | Naproxen (I)                     | Liquid extraction  | [341] |
|        |        |     |        |     |   | CE              | Forensic              | DNA                              | Experimental details for sample extraction   | [342] |

Table 7. Continued

| Coupling mode |    |    |        | CFS |   | Electrophoretic | Sample                 | Analytes                          | Remarks  | Ref.  |
|---------------|----|----|--------|-----|---|-----------------|------------------------|-----------------------------------|--|-------|
| OFF           | IN | ON | OFF/ON | NC  | С | mode            |                        |                                   |  |       |
| x             |    |    |        | X   |   | CE              | _                      | Sulfonated metallophtalocyanines  | Soxhlet extraction                                   | [343  |
| K             |    |    |        | X   |   | CE              | Wheat flour            | Gliadins                          | Sample was extracted with buffer solution            | [344  |
| (             |    |    |        | X   |   | CE              | Soil                   | Platinum                          | Extraction of platinum after static extraction       | [345  |
|               |    |    |        | X   |   | CE              | DNA                    | Nucleic acids/deoxyribo adducts   | Review (extraction as sample preparation)            | [346  |
| (             |    |    |        | X   |   | CE              | Fish and crab meat     | Methylmercury                     | Pretreatment with different organic solvent          | [34]  |
| (             |    |    |        | X   |   | CE              | Environmental samples  | Chromate/aromatic hydrocarbons    | Extraction media buffer                              | [348  |
| K             |    |    |        | X   |   | CE              | Hair                   | Amphetamine derivatives           | Liquid-liquid extraction                             | [349  |
| K             |    |    |        | X   |   | CE              | Cocaine                | Isomeric truxillines              | Liquid-liquid extraction                             | [350  |
| K             |    |    |        | X   |   | CE              | Enantiomers mianserin  | Blood plasma                      | Liquid-liquid extraction                             | [35]  |
| ζ             |    |    |        | X   |   | CE              | Inks                   | _                                 | Optimisation of the extraction conditions            | [352  |
| ζ             |    |    |        | X   |   | CE              | Plasma                 | Phenylacetic acid                 | Simultaneous extraction and preconcentration         | [353  |
| (             |    |    |        | X   |   | CE              | Pharmaceutical         | Oxytetracycline (I)               | Extraction with N-methylformamide (15 min)           | [354  |
| X             |    |    |        | X   |   | CE              | Body fluids            | Toxic drugs                       | Liquid-liquid extraction                             | [355  |
| X             |    |    |        | X   |   | CZE             | Wool                   | Dyes                              | Dyes were extracted from wool                        | [356  |
| X             |    |    |        | X   |   | CZE             | Urine                  | Methylenedioxymethamphetamine     | Hydrolysis was carried out prior extraction          | [357  |
| X             |    |    |        | X   |   | CE              | Water                  | Whitening agent (tinopal)         | Styrene-divinylbenzene extraction discs              | [358  |
| X             |    |    |        | X   |   | CE              | Blood/serum            | Flunixin                          | Extraction of flunixin on C <sub>18</sub> cartridges | [359  |
| ζ.            |    |    |        | X   |   | CGE             | Plasma/urine           | Phosphorothioate oligonucleotides | Extraction with phenol–CHCl <sub>3</sub>             | [360  |
|               |    |    |        | x   |   | MEKC            | Algal scum             | Toxins                            | 10 ml SFE vessel                                     | [36]  |
| ζ.            |    |    |        | X   |   | CE              | -                      | Enantiomers of propranolol        | Soxhlet extraction                                   | [362  |
| ζ.            |    |    |        | x   |   | CE              | Urine                  | Dimethindane                      | Extraction with cyclohexane–ethyl acetate            | [363  |
| ζ.            |    |    |        | X   |   | MEKC            | Biological fluids      | Paclitaxel                        | Extraction with cyclonexane—ethyl acetate            | [364  |
| X.            |    |    |        | X   |   | CE              | Diological fluids      | Nucleic acids                     | Genomic DNA was extracted from lung                  | [365  |
|               |    |    |        |     |   | MEKC            | Human urine            | Antidepressants                   | Extraction was performed with hexane                 | [366  |
| X             |    |    |        | X   |   | MEKC            | Cozaar tablets         | Losartan potassium drug           | •  | [367  |
| ,             |    |    |        | X   |   |                 |                        |                                   | Robotic extraction with buffer                       | -     |
|               |    |    |        | X   |   | CE              | Freeze-dried tuna      | Methylmercury                     | Sample was extracted with acetone–water              | [368  |
| X             |    |    |        | X   |   | CE              | Environmental samples  | Methylmercury                     | Extracted with acetone and toluene                   | [369  |
| · ·           |    |    |        | X   |   | CE              | Dissolution test       | Clenbuterol and levothyroxine     | Extraction discs                                     | [370  |
| K.            |    |    |        | X   |   | CE              | DNA                    | Benzo[a]pyrene diol epoxide       | Unconsumed diol was removed by extraction            | [37]  |
|               |    |    |        |     |   | CE NEWS         | -                      | -                                 | Review of on-line sample extraction                  | [372  |
| X             |    |    |        | X   |   | CE-MEKC         | Soil samples           | Inorganic ions                    | Description of the extraction of analytes            | [373  |
| X             |    |    |        | X   |   | CE              | Amniotic fluid         | DNA markers (Down's syndrome)     | Phenol-CHCl <sub>3</sub> extraction                  | [374  |
| X             |    |    |        | X   |   | CZE             | DNA                    | Styrene oxide adducts             | Unreacted products removed by extraction             | [375  |
| X             |    |    |        | X   |   | CZE             | Cotton and wool fibres | Black dyes                        | Extraction with NaOH (25 min+100°C)                  | [376  |
| X             |    |    |        | X   |   | CZE             | Grass pea              | 3-N-Oxalyl-diaminopropanoic acid  | Extraction twice by shaking (45 min)                 | [377  |
| K             |    |    |        | X   |   | CZE             | Natural products       | Nitrate, nitrite, amino acids     | Extraction of anti-cancer natural products           | [378  |
| ζ.            |    |    |        | X   |   | MEKC-CE         | Urine                  | Drugs of abuse                    | Extraction scheme                                    | [379  |
| K             |    |    |        | X   |   | CE              | Chinese medicine       | Six bioactive components          | Sample was extracted with methanol                   | [380  |
| K             |    |    |        | X   |   | CZE             | Milk                   | Chloroamphenicol                  | Extraction with ethyl acetate                        | [38]  |
| ζ.            |    |    |        | X   |   | CE              | Urine/plasma           | 7-Hydroxycoumarin                 | Extraction with diethyl ether (10 min)               | [382  |
| K             |    |    |        | X   |   | CE              | Organic solvent        | Cationic and anionic              | Ions were concentrated by extraction                 | [383  |
| Κ             |    |    |        | X   |   | MEKC            | Water                  | Monosulfonated dyes               | Dyes were isolated by extraction                     | [384  |
| Κ             |    |    |        | X   |   | MECC            | Pharmaceutical         | Anabolic steroids                 | A one step quantitative extraction procedure         | [386  |
| K             |    |    |        | X   |   | MEKC            | Human serum            | Drugs                             | Solvent extraction procedures                        | [388] |
| X             |    |    |        | X   |   | CE              | Hair                   | Cocaine                           | Extraction with Toxi-Tubes A                         | [389  |
| K             |    |    |        | X   |   | CE              | Rat hair               | Amino acids                       | Extraction with CHCl3-propan-2-ol                    | [390  |
| K             |    |    |        | X   |   | MEKC            | Serum                  | Amino acids                       | Extraction with pentane                              | [39]  |
| Κ             |    |    |        | X   |   | CE-ITP          | _                      | Anionic, cationic and drugs       | Liquid-liquid extraction                             | [392  |
| K             |    |    |        | X   |   | CE              | Foods                  | Glycine, 5-hydroxymethylfurfural  | Aldehydes were removed by extraction                 | [393  |
| ĸ             |    |    |        | x   |   | CZE-MEKC        | Soil/water             | Aromatic organic acids            | Water was extracted with extraction disks            | [394  |
|               |    |    |        |     |   | CZE             | Orange juice           | Ascorbic and dehydroascorbic acid | Extraction described in the article                  | [395  |

Table 7. Continued

| Coupli | Coupling mode CFS |    |        |    |   | Electrophoretic | Sample             | Analytes                 | Remarks                                       | Ref.  |  |
|--------|-------------------|----|--------|----|---|-----------------|--------------------|--------------------------|---|-------|--|
| OFF    | IN                | ON | OFF/ON | NC | С | mode            |                    |                          |   |       |  |
| x      |                   |    |        | X  |   | CZE-MEKC        | Human urine        | Morphine 6-glucuronide   | Extraction improved the LOD                   | [396] |  |
|        |                   |    |        |    |   | MEKC            | Foodstuffs         | Mycotoxins               | Review (emphasis of sampling, extraction)     | [397] |  |
| X      |                   |    |        | X  |   | CE              | Dystrophin gene    | Nucleic acids            | The extraction was with phenol-chloroform     | [398] |  |
| X      |                   |    |        | X  |   | CZE             | Sugar cane         | Flavonoids               | Acetonitrile-water extraction prior to CZE    | [399] |  |
| X      |                   |    |        | X  |   | CZE             | Blood/serum        | Pentobarbitone           | Extraction gave an increase in sensitivity    | [400] |  |
|        |                   |    |        |    |   | MEKC            | Biological fluids  | Drugs                    | Overview                                      | [401] |  |
| X      |                   |    |        | X  |   | CZE             | Rat urine          | Cimetidine               | Extraction with ethyl acetate-light petroleum | [402] |  |
| X      |                   |    |        | X  |   | CE              | Coptidis Rhizoma   | Quaternary alkaloids     | Extraction was repeated three times           | [403] |  |
| X      |                   |    |        | X  |   | CE              | Blood/plasma       | Cicletanine              | Extraction was with diethyl ether             | [404] |  |
| X      |                   |    |        | X  |   | MEKC            | Residues           | Gunshot                  | Extraction with ethanol                       | [405] |  |
| X      |                   |    |        | X  |   | MEKC            | Human serum/plasma | Thiopental (thiopentone) | Liquid-liquid extraction                      | [406] |  |
| X      |                   |    |        | X  |   | CE-MEKC         | Human serum/urine  | Barbiturates             | Extraction with cartridges                    | [407] |  |
| X      |                   |    |        | X  |   | CZE             | Organic soils      | Caesium                  | Extraction with NaOH under $N_2$              | [455] |  |

CFS: Continuous flow system; NC: no chromatography; C: chromatography; CZE: capillary zone electrophoresis; CE: capillary electrophoresis; MEKC: micellar electrokinetic chromatography; ITP: isotachophoresis; SPE: solid-phase extraction; LOD: limit of detection; FI: flow injection; CGE: capillary gel electrophoresis; HIV: human immunodeficiency virus; SFE: supercritical fluid extraction.

sample solutions were injected into the ITP system comprising a PTFE separation capillary and a fused-silica detection capillary with on-line UV detection. The ITP and CE systems were coupled via a Plexiglass interface block.

In 1996, Sadecka and Polonsky [52] carried out the determination of some cardiovascular drugs in serum and urine by capillary ITP. The analytical column was connected to a preseparation column. Amiloride and β-blockers were separated by cationic isotachophoresis in a 10 mM sodium morpholinoethanesulfonate buffer (pH 5.5)–5 mM glutamic acid system. Frusemide was separated in the anionic electrolyte system 10 mM histidine hydrochloride buffer (pH 6.2)–morpholinopropanesulfonic acid.

### 12. High-performance liquid chromatography coupled to capillary electrophoresis

The coupling of LC and CE was described by Bushey and Jorgenson [53]. As CE operates under fundamentally different separation mechanisms, the combination with LC represents a true orthogonal system. A reversed-phase LC (RPLC) system was used in the first dimension, and eluting fractions were introduced and further separated on a CE system, which was used to separate peptide standards

and fluorescently labelled peptide fragments from a tryptic digest of ovalbumin. The coupling HPLC–CE is not yet widely used, 13 references are included in Table 9.

Larman Jr. et al. [54] developed a two-dimensional separations of peptides and proteins by comprehensive LC-CE. Tryptic digest of horse heart cytochrome c, labelled with fluorescein isothiocyanate, was subjected to two-dimensional RPLC and fast CZE (FCZE). Eluates passed through an untreated fused-silica capillary and interface tee which led to waste (to lower the pressure) and into the second dimension FCZE system with sampling controlled by an argon ion laser gating beam.

Moore Jr. and Jorgenson [55] developed a comprehensive three-dimensional separation of peptides using size-exclusion chromatography-RPLC-optically gated CZE. The LC column was linked to the top of a vertical fused-silica CZE capillary, with a high-speed optical gating injection system.

A transparent flow gating interface for the coupling of microcolumn LC with CZE in a comprehensive two-dimensional system was developed by Hooker and Jorgenson [56]. The new interface was based on the original flow gated design developed in their laboratory but is now made from a clear plastic which allows for the direct observation and routine manipulation of the micro-HPLC and CZE capillaries. To evaluate the reproducibility of the inter-

Table 8
Coupling continuous isotachophoresis systems to capillary electrophoresis

| Coup | oling  | mode |        | CFS  | Electrophoretic | Sample                                   | Analytes                          | Remarks  | Ref |
|------|--------|------|--------|------|-----------------|--|-----------------------------------|--|-----|
| )FF  | IN     | ON   | OFF/ON | NC C | mode            |  |                                   |  |     |
|      |        | Х    |        | х    | CE              | Proteins                                 | Derivatives of amino acids        | ITP as concentration pretreatment technique in CE            | [4  |
|      |        | X    |        | X    | CZE             | _  | Eight proteins                    | CZE with on-column transient ITP preconcentration            | [49 |
|      |        | X    |        | X    | ITP-CZE         | _  | Protein                           | On-column transient and coupled-column ITP                   | [5  |
|      |        | X    |        | X    | CE              | _  | Derivative of angiotensin III     | Analyte focusing in CE using on-line ITP                     | [5  |
|      |        | X    |        | X    | ITP             | Serum/urine                              | Cardiovascular drugs              | Analytical and preseparation column connected                | [5: |
|      |        |      |        |      | CE              | Aqueous samples                          | Environmental pollutants          | Overview   | [1  |
|      |        |      |        |      | CE              | _  | -                                 | Membrane preconcentration system                             | [1: |
|      | X      |      |        | x    | CZE             | Urine                                    | Adenosine                         | CZE with oncolumn ITP preconcentration                       | [2: |
|      | Х      |      |        | X    | CZE             | Blood plasma                             | Heterocyclic peptides             | Improved sensitivity by on-line ITP preconcentration-CZE     | [2: |
|      | X      |      |        | X    | CE              | -  | Peptides                          | On-line transient ITP  | [2  |
|      | Х      |      |        | X    | CE              | Urine                                    | Peptides and proteins             | Analytes can be submitted to stacking or ITP                 | [2  |
|      |        |      |        | X    | CE              | -  | Peptides and proteins             | ITP conditions for membrane preconcentration–CE              | [2  |
|      | X      |      |        | X    | CZE             |  | Peptide mixtures                  | CE–MS in conjunction with transient ITP                      | [3  |
|      | X      |      |        | X    | CZE             | Antimuscarinic drugs                     | Anionic and cationic              | ITP preconcentration for CZE in a single capillary           | [3  |
|      |        |      |        | X    | CZE-ITP         | Human erythrocytes                       | Adenosine deaminase activity      | On-column capillary ITP–CZE                                  | [4] |
|      | X<br>X |      |        | X    | CE-III          | Nuclear fission product                  | Rare-earth metals                 |  |     |
|      | Х      |      |        |      |                 | Nuclear fission product                  |                                   | ITP permits stacking of large injection volumes              | [4] |
|      |        | X    |        | X    | CZE             | -<br>-                                   | Sulfanilic and hippuric acids     | CZE with on-line ITP sample pretreatment                     | [4] |
|      |        | X    |        | X    | CZE             | Antimuscarine drugs                      | -                                 | Counter-flow ITP-CZE   | [4  |
|      | X      |      |        | X    | CZE-ITP         | Pseudomonas aeruginosa                   | Oligosaccharides                  | Transient ITP preconcentration coupled to CZE                | [4  |
|      | X      |      |        | X    | CE              | Biological cells                         | Antisense oligonucleotides        | On-capillary ITP and CE polymer sieving                      | [4  |
|      | X      |      |        | X    | CE              | _  | Cholinesterase inhibitor          | ITP was used to concentrate before CZE                       | [4  |
|      | X      |      |        | X    | CE              | Fission products                         | 137Cs, 152Eu and lanthanide       | ITP enabled large volume samples to be analysed              | [4  |
|      |        | X    |        | X    | CE              | Blood serum                              | Hippuric acid                     | On-line ITP-CZE  | [4  |
|      | X      |      |        | X    | CE              | Drugs                                    | Amitriptyline and metoprolol      | ITP preconcentration with CZE in a single capillary          | [4  |
|      |        |      |        |      | CE              | -  | -                                 | Review (combined technique of capillary ITP-CZE)             | [4  |
|      |        | X    |        | X    | CE              | Natural, mineral water                   | EDTA-iron(III) complex            | On-line coupling of capillary ITP and CZE                    | [4: |
|      |        |      |        | X    | CZE             | Water                                    | Nitrophenols                      | Off-line isotachophoretic sample pretreatment                | [42 |
|      |        | X    |        | X    | CZE             | Mayonnaise                               | EDTA                              | On-line coupled capillary ITP-CZE                            | [42 |
|      |        |      |        | X    | CE              | -  | Oligonucleotides                  | On-column transient capillary ITP and CE                     | [42 |
|      | X      |      |        | X    | CZE             | -  | Nitrophenols                      | To enhance the sample load capacity: ITP-CZE                 | [42 |
|      |        |      |        | X    | CZE             | Basic proteins                           | Basic peptides                    | On-line ITP-CZE with hydrodynamic counterflow                | [4: |
|      | X      |      |        | X    | CGE             | X174/HaeIII and PCR<br>product of 118 bp | DNA                               | ITP preconcentration in capillary gel electrophoresis        | [42 |
|      |        | X    |        | X    | CZE             | -  | _                                 | ITP (preseparation and concentration) and CZE                | [4: |
|      | X      |      |        | X    | CZE             | _  | _                                 | Review (single capillary coupled ITP-CZE techniques)         | [4  |
|      |        | X    |        | X    | CZE             | β-Blocker drugs                          | _                                 | Combined LLE-ITP for loadability in CZE                      | [4: |
|      | x      |      |        | X    | CZE             | -  | Inorganic anions                  | On-line coupled capillary ITP-CZE                            | [4  |
|      | X      |      |        | X    | CZE             | _  | Neostigmine and others            | Combined LLE and ITP as fast on-line focusing step in CE     |     |
|      | Х      |      |        | X    | CZE             | Calf urine                               | β-Agonists                        | On-capillary ITP for loadability enhancement in CZE          | [4: |
|      | X      |      |        | X    | CZE             | -  | Neostigmine/propantheline         | In-line ITP focusing of large injection volumes for CZE      | [4: |
|      | A      | X    |        | X    | CZE             | _  | Amino acids                       | Automated on-capillary ITP followed by CZE.                  | [4  |
|      | v      | А    |        | X    | CZE             | _  | Brilliant acid green              | Correlation between velocity and current in ITP-CZE          | [4  |
|      | X<br>X |      |        | X    | CZE             | Tap or lake water                        | Paraquat and diquat               | ITP sample pretreatment and determination by CZE             | [4  |
|      |        |      |        |      | CZE             | -  | Enzyme inhibitor                  |  | [4  |
|      | X      |      |        | X    |                 | Biological samples                       | Elizythe illiliottor              | On-column ITP focusing in CZE                                | -   |
|      | X      |      |        | X    | CE              | Protein                                  | -<br>C-16:1/2 5 di:1:1            | On-column transient capillary ITP preconcentration in CE     | [4: |
|      |        | X    |        | X    | CZE             | Urine                                    | Sulfanilate/3,5-dinitrosalicylate | CZE of mixtures with on-line ITP sample pretreatment         | [4  |
|      |        | X    |        | X    | CZE             | -  |                                   | Options in electrolyte systems for combined ITP-CZE          | [4  |
|      |        | X    |        | X    | CZE             | Dilute sample                            | p- and m-chlorobenzoic acids      | Analysis by CZE with ITP preconcentration                    | [4  |
|      |        | X    |        | X    | CE              |  |                                   | On-line coupling of capillary ITP-CE                         | [4  |
|      | X      |      |        | X    | CZE             | Feedstuffs                               | Halofuginone                      | Combination of ITP-CZE in a column-switching system          | [4  |
|      |        | X    |        | X    | CZE             | Blood                                    | Anionic/cationic, thiamine        | ITP preconcentration for enhancement of detectability in CZE |     |
|      | X      |      |        | X    | CZE             | -  | Cytochrome                        | CZE of dilute samples with ITP preconcentration              | [4  |
|      |        | X    |        | X    | CZE             | -  | Nitrophenols/labelled amino acids | On-line coupling of capillary ITP with CZE                   | [4  |
|      | X      |      |        | X    | CZE             | _  | Polypeptides and proteins         | CZE and ITP-mass spectrometry                                | [4  |
|      |        |      |        |      | ITP             | _  |                                   | Review   | [4  |
|      | X      |      |        | X    | ITP             | Water                                    | Permethrin and tetramethrin       | Double extraction  | [4: |
|      | X      |      |        | X    | ITP             | Non-aqueous                              | Cationic metal chelates           | Capillary-tube ITP-solvent extraction                        | [4: |

CFS: Continuous flow system; NC: no chromatography; C: chromatography; CZE: capillary zone electrophoresis; CE: capillary electrophoresis; ITP: isotacophoresis; LLE: Liquid-liquid extraction.

Table 9
Coupling continuous HPLC systems to capillary electrophoresis

| Couplin | g mode |    |        | CFS |   | Electrophoretic | Sample                        | Analytes              | Remarks  | Ref.  |
|---------|--------|----|--------|-----|---|-----------------|-------------------------------|-----------------------|--|-------|
| OFF     | IN     | ON | OFF/ON | NC  | С | mode            |                               |                       |  |       |
|         |        |    |        |     | х | CE              | -                             | -                     | Review (multi-dimensional separation systems)                    | [19]  |
|         |        | X  |        |     | x | CE              | Ovalbumin                     | Peptides              | 2D HPLC-CE   | [53]  |
|         |        | X  |        |     | X | CZE             | Horse heart cytochrome        | Peptides and proteins | 2D separations by LC-CE  | [54]  |
|         |        | X  |        |     | x | CE              | Hen ovalbumin                 | Peptides              | Three-dimensional separation: SEC-LC-CE                          | [55]  |
|         |        | x  |        |     | X | CZE             | Urine                         | Amino acids           | Coupling of microcolumn LC with CZE                              | [56]  |
|         |        | X  |        |     | X | CE              | -                             | Benzoic acids         | On-line chromatographic pretreatment of samples for CE           | [57]  |
| x       |        |    |        |     | x | CZE             | Plasma                        | Drugs                 | Micro-CLC as an interface between SLM extraction and CZE         | [287] |
| x       |        |    |        |     | x | CE              | Endoproteinase Lys-C          | Thirty glycoforms     | CE resolved fractions from the LC                                | [408] |
|         |        |    |        |     | x | CE              | _                             | Metallothioneins      | Review (coupling of separation techniques)                       | [409] |
|         |        |    |        |     | x | ITP             | Soil                          | Herbicide asulam      | HPLC as a sample prep. method                                    | [452] |
|         |        | x  |        |     | X | CZE             | Proteins                      | _                     | Transverse flow gating interface for the coupling of LC-CZE      | [453] |
|         |        | X  |        |     | x | CZE             | _                             | Peptides              | Analytes were separated by 2D HPLC and CZE                       | [454] |
|         |        | x  |        |     | x | CE              | _                             | PAHs                  | Application of 2D statistical theory                             | [456] |
|         |        | X  |        |     | X | CEC             | Energetic materials standards | Nitro compounds       | Simple interface for gradient CEC, supplies mobile phase from LC | [457] |

CFS: Continuous flow system; NC: no chromatography; C: chromatography; CZE: capillary zone electrophoresis; CE: capillary electrophoresis; SPME: solid-phase microextraction; MEKC: micellar electrokinetic chromatography; CGE: capillary gel electrophoresis; CEC: capillary electrochromatography; ITP: isotachophoresis; PAHs: polynuclear aromatic hydrocarbons; SEC: size-exclusion chromatography; CLC: column liquid chromatography; SLM: supported liquid membrane; 2D: two-dimensional.

face, 400 consecutive CZE separations of a mixture of fluorescein 5-isothiocyonate (FITC)-labelled phenylalanine and glutamic acid were performed.

The direct injection of samples with high salt concentrations in CE results in peak splitting and/or serious band broadening. These problems were not encountered when using a liquid chromatographytype of sample pretreatment coupled on-line with a CE system. To demonstrate the feasibility of this approach, the separation of three model compounds (benzoates) in water containing up to 400 mM of sodium chloride was studied [57].

#### 13. Conclusions

CE has many weak points that can be attributed to the sample preparation step. As has been described in this review, a lot of ingenious CFSs have been designed to solve a variety of problems encountered in CE analysis. Even though it is not easy to predict the future of all those smart inventions (will they survive and become widely accepted or not) they all enrich the field of CE. In summary, the different techniques presented in this review can overcome the current limitations of the LODs in CE and reduce the need for exhaustive off-line sample preparation prior to analysis by CE. This technique is slowly gaining popularity as a tool with great potential for routine analysis in the laboratory.

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