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1. Introduction

Mass spectrometry (MS) has become a powerful analytical tool for qualitative and quantitative applications, providing information about the structure and purity of compounds, and also about the chemical composition of complex samples.

The most recent applications of mass spectrometry are oriented towards biochemical applications such as proteome, metabolome and drug discovery. During the last decade, mass spectrometry has progressed rapidly and an evolution has been observed in the type of applications, software and equipments. Atmospheric pressure ionization sources are now used, an analyser based on a new concept (the orbitrap) was recently developed, existing ones were modified, and new hybrid instruments were developed using combinations of different analysers, depending on applications. One of the major trends was the transition to high resolution/accurate mass analysis, made routine by new MS instruments. The use of separation techniques as gas chromatography (GC), liquid chromatography (LC) and capillary electrophoresis (CE) coupled with mass spectrometry and tandem mass spectrometry, expanded the interest in this methodology.

In this chapter are presented general aspects related with characteristics of mass spectrometry equipments. The contribution of this technique to new discoveries concerning one of the major infectious diseases in man, malaria, is also discussed.

2. Basics of mass spectrometry

Mass spectrometry is a technique used to analyse from small inorganic molecules to biological macromolecules and relies on the formation of gas-phase ions (positively or negatively
In order to achieve this state, the sample must be volatilized and this may become a problem to biological samples, as biomolecules have usually high molecular mass and high polarity, factors that limit their volatility.

All mass spectrometers share common components as an ionization source, a mass analyser and a detector (Fig. 1). As there are available equipments with different specifications, even from the same supplier, it is necessary to choose carefully the most adequate equipment for each type of application.

Figure 1. Basic components in a mass spectrometer. ESI, electrospray ionization, APCI, atmospheric-pressure chemical ionization; MALDI, Matrix-Assisted Laser Desorption ionization; DESI: Desorption Electrospray Ionization ; DART: Direct Analysis in Real Time; FT-ICR: Fourier transform ion cyclotron resonance (adapted from Glish & Vachet, 2003)

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Different ionization techniques may be used in mass spectrometry equipments, depending on the need of molecule disruption for the induction of ion formation. These techniques may perform strong and soft ionization processes. Soft ionization methods, like fast atomic bombardment (FAB), liquid secondary ion mass spectrometry (LSIMS), matrix-assisted laser desorption ionization (MALDI) and electrospray ionization (ESI) allow the detection of molecular ions and are more suitable for the analysis of biomolecules and non-purified analytes. The ions are generated by protonation, deprotonation or formation of adducts. In table 1 are summarized some of the main characteristics of the most used ionization methods.

The use of FAB is useful to assign the molecular ion peaks. The sample is dissolved in a suitable liquid matrix, with low vapour pressure (e.g. diethanolamine, triethanolamine, glycerol, thioglycerol or 3-nitrobenzyl alcohol), inserted into the mass spectrometer and bombarded with high energy argon or xenon atoms, providing efficient means to analyze polar, ionic, thermally labile, energetically labile, and high molecular mass compounds (El-Aneed et al., 2009).
In MALDI, ions are produced by pulsed-laser irradiation (e.g. nitrogen lasers) of the sample co-crystallized with an organic matrix (e.g. gentisic, sinapic or ferulic acid) and operating in the vacuum or more recently, at atmospheric pressure. MALDI ionization uses a low amount of sample but low molecular mass molecules (below 500 Da) are difficult to analyse, due to strong interferences of the organic matrix ions.

The development of other soft ionization techniques has become crucial to the analysis of biomolecules dissolved in a mixture of water and a volatile organic solvent (e.g. methanol, acetonitrile). Techniques as ESI and MALDI make MS methodologies versatile as both techniques accomplish the conversion into gas-phase ions of non-volatile and thermally labile large molecules, allowing the study of biological compounds. Both techniques produce protonated peptide and protein ions, deprotonated deoxyribonucleic acids (DNA) and ribonucleic acids (RNA). Some reviews have been published on the covalent and non-convalent interactions between drug molecules with DNA and RNA, protein and enzyme targets for drug action and toxicity (Feng, 2004). When using ESI, proteins are ionized as they have several sites of protonation or deprotonation, and this multiple charging enables mass spectrometers with limited m/z ranges to analyse higher molecular mass molecules. However, ion suppression may occur when solutions contain high concentrations of salt, or when the target analytes are present in low concentration in matrices with high content of other analytes. Strategies based on the type of analyte, ionization reaction, ionization efficiency, analyte solution composition and pH, have been described for producing positive or negative ion modes when operating with an ESI source (Feng, 2004). APCI, is less susceptible to matrix interferences from salts, and is used for monitoring weakly polar compounds. However, labile

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Table 1. Characteristics of the most used ionization sources. FAB: Fast Atomic Bombardment; MALDI: Matrix Assisted Laser Desorption Ionization; ESI: Electrospray Ionization; DESI: Desorption Electrospray Ionization; DART: Direct Analysis in Real Time
Compounds can be thermally decomposed, and due to its high sensitivity, the solvents used with this technique must have higher purity.

DART and DESI are well established open-air ionization techniques, as no sample preparation is required, making these techniques suitable for screening a large number of samples (Fernández et al., 2006). The DART ion source produces a heated stream of protonated reactant ions and the analytes in the sample are ionized, producing protonated molecules $[M+H]^+$ or deprotonated molecules $[M-H]^-$ in the open air of the laboratory environment, making possible the analysis of organic compounds directly, in real time, without time-consuming analytical protocols and destruction of the sample. The method may detect concentrations of analytes as low as femtomole (Arnaud, 2007). Due to these characteristics, DART has become an ionization method useful for rapid screening of pharmaceutical products. In DESI analysis, a high-speed charged liquid spray is directed to the sample (Takats et al., 2005). The DESI spray dissolves the material from the sample and the charged droplets are sampled downstream by a mass spectrometer. Desolvation and evaporation from these droplets creates ions that will generate a mass spectrum of the sample components.

Following the ionization process, the selected ions are extracted, accelerated, and analyzed. A mass analyser is characterized by its mass range limit, analysis speed, transmission, mass accuracy and resolution, expressed as full width at half maximum (FWHM). The most used analysers and their characteristics are summarized in Table 2.

Quadrupoles are widely used mass analysers, where ions are separated according to the stability of their trajectories in the oscillating electric fields applied between the four parallel rods. The QTOF, a hybrid quadrupole time of flight mass spectrometer, is a high-resolution mass spectrometer with MS/MS capability, and has been often used in drug studies (Nyunt et al., 2005). FT-ICR is also a high resolution and high mass accuracy analyser that enables the study of the binding of ligands (drugs) to RNA targets (Hofstadler et al., 1999; Masselon et al., 2000). The Orbitrap mass analyzer employs electrostatic trapping and it bears similarities to FT-ICR as both belong to the same Fourier Transform MS (FTMS) family of instruments. Orbitrap mass spectrometry is expected to provide maximum resolving powers of 100,000–200,000. A modified Orbitrap instrument has shown that this technology is capable of a resolution of 1,000,000 for m/z < 300–400, which makes it compatible to be used with chromatographic separation techniques (Denisov et al., 2012).

Tandem mass spectrometry (MS²) development was crucial for the structural analysis of compounds. In tandem experiments, a molecular ion is selectively isolated and fragmented in a controlled environment. With this type of analysers, it is possible to perform different types of experiments (e.g. parent scan, daughter scan, neutral loss) and data obtained, will allow to identify or quantify the analytes, even in complex matrices (e.g. natural product extracts and biological fluids). Multiple Reaction Monitoring (MRM), has become an important tool, used for quantification purposes, allowing an increment of methods specificity and sensitivity.

Finally the signal obtained in the detector will produce a mass spectrum, the x-coordinate represents m/z values and the y-axis indicates total ion counts.
Liquid chromatography coupled to mass spectrometry (Triple Quadrupole TQ, QTOF, Linear ionTrap and Linear QTRAP analyzers) is today a well established methodology used due to its high sensitivity, speed, selectivity, versatility and ease of automation. Recently the advantages of using mass spectrometry in comprehensive liquid chromatography (LC X LC system) have been discussed and different applications have been described for pharmaceutical compounds (Donato et al., 2012).

3. An overview of malaria

Malaria is caused by *Plasmodium* parasites, which are transmitted through the bite of an infected female *Anopheles* mosquito, and remains one of the major infectious disease in man.

Five species from the genus *Plasmodium* namely: *P. falciparum*, *P. vivax*, *P. ovale*, *P. malariae*, and *P. knowlesi* cause infection in humans. Of these, *P. falciparum* and *P. vivax* account for more than 95% of malaria cases in the world, with *P. falciparum*, being responsible for most of the deaths caused by malaria every year. The species of human malaria differ in the periodicity of their
life cycle, as well as in the outcomes of the disease. Generally clinical manifestations can include fever, chills, prostration and anemia. Severe disease can include delirium, metabolic acidosis, cerebral malaria and multi-organ system failure, coma and death may ensue. (Kantele & Jokiranta, 2011)

The World Health Organization (WHO) estimated 225 million cases of malaria and about 800,000 deaths worldwide in 2010 (WHO, 2010). Due to the rapid evolution and spread of multi-resistant parasites to the current antimalarial drugs, both, chemotherapy and prophylaxis are at risk of impairment. Malaria is most prevalent in developing countries of tropical areas such as sub-Saharan Africa, East Asia and South America (Rodrigues et al., 2010; Eisenstein, 2012).

3.1. Life cycle of the malaria parasite

The malaria parasite exhibits a complex life cycle (Fig. 2) involving an insect vector (mosquito) and a vertebrate host (human). It includes an asexual cycle in humans, encompassing an asymptomatic liver-stage, a symptomatic blood-stage and a sexual cycle in a mosquito.

The liver or hepatic stage (A) is initiated when sporozoites injected through the bite of a mosquito travel to the liver and infect hepatocytes, where a clinically silent asexual multiplication takes place, generating thousands of merozoites. The release of merozoites into the bloodstream (B) marks the beginning of the erythrocytic stage of infection (C), during which parasites infect red blood cells, undergo repeated asexual replication cycles, and give rise to clinical illness. Some merozoites differentiate into gametocytes (D) that can be taken up by the mosquito during a posterior blood meal. Within the mosquito, gametocytes undergo a sexual development to form sporozoites that migrate to the salivary glands and can infect another host through another bite (E).

Figure 2. Plasmodium life cycle
3.2. Agents with antimalarial activity

The public health problem of malaria has been addressed by different approaches (Biot et al., 2012):

- use of insecticides to control the mosquito vector, but mosquitoes are developing resistance to insecticides;
- vaccines, but in spite of all the efforts there is yet not available a vaccine that effectively targets the parasite;
- chemotherapy to control malaria has relied mainly on a restricted number of chemically related drugs belonging to either the quinoline or the antifolate groups.

Increasing resistance of *P. falciparum* to the commonly used drugs is recognized as one of the major problems in eradication of the disease. The severe malaria situation underscores the continuing need of research for new classes of antimalarial agents with new mechanisms of action or re-utilization of the existing drugs with new types of therapies. The existing drug armamentarium is insufficient to answer the call for malaria eradication. The first line of treatment for malaria currently relies on a single class, the artemisinins. To overcome this problem scientists are exploring many approaches, targeting different stages of the parasite life cycle, to find agents that will prevent, cure or eliminate malaria. (Hobbs, C. & Duffy, P., 2011)

Many antimalarial agents contain a 4-aminoquinoline, 8-aminoquinoline or quinolone methanol scaffolds (Rosenthal, 2001). Chloroquine and amodiaquine are 4-aminoquinolines used to treat and prevent malaria, while primaquine is the single 8-aminoquinoline clinically approved to treat relapsing malaria caused by *P. vivax*. Mefloquine (Fig. 3) is a quinoline methanol antimalarial structurally similar to quinine, the first pure substance used to treat malaria and extracted from the bark of the cinchona tree. Other relevant classes of antimalarial agents include the antifolates (e.g. pyrimethamine and proguanyl), phenanthrene methanols (e.g. halofantrine), and naphthoquinones (e.g. atovaquone). More recently, artemisinin (Fig. 3), a sesquiterpene lactone isolated from the *Artemisia annua* chinese herb, and its analogues were a major breakthrough in malaria chemotherapy because they produce a very rapid therapeutic response, particularly against multidrug-resistant *Plasmodium falciparum* malaria.

With exception of primaquine, (Vale et al., 2009) most available antimalarials are active against the blood stage of the disease. However to achieve the eradication goal, new compounds with new modes of action are needed to block parasite transmission and eliminate the asymptomatic and latent hepatic forms. (Rodrigues et al., 2012) A strategy used to address this major goal is to combine two chemotypes - each one targeting a specific stage of the parasite’s life cycle in a single chemical entity, to develop effective hybrid antimalarials capable of killing both the blood and liver-stage parasites with identical efficacy. (Capela et al., 2011)
4. Study of antimalarials agents by mass spectrometry

Structural and stability information is fundamental for any drug study, including antimalarials. In fig. 3 are presented currently available antimalarial drugs. Due to the rapid emergence and spread of resistant parasites to well-established antimalarial drugs, there is an urgent need for novel drugs. Studies performed on new antimalarial compounds using mass spectrometry are scarce but are useful for the elucidation of structures, also for prediction of compound stability and properties, isomer characterization, and detection of counterfeit products. Furthermore, studies using this technique coupled to chromatographic methods have been conducted for the evaluation of pharmacokinetics, metabolite identification, and detection of impurities.

4.1. Structural elucidation

One of the main applications of mass spectrometry is the structural elucidation of molecules. Based on the molecular ion peaks and their fragmentation patterns, the structure of compounds can then be proposed.

Among the different equipments that can be used for these purposes, FAB ion sources are frequently described. Applications can be found in the study of the oxidation products of primaquine, 5,5-di-[6-methoxy-8-(4-amino-1-butyl amino)] quinoline (PI), 6-methoxy-5,8-di-[4-amino-1-methyl butyl amino] quinoline (PII) and 5,5-di-[7-hydroxy-6-methoxy-8(4-amino-1-methyl butylamino)] quinoline (PIII) (Fig. 4) (Sinha & Dua 2004). The mass spectrum of PI with molecular formula $C_{30}H_{40}N_6O_2$ presents the molecular ion at $m/z$ 517 confirming the molecular mass of the compound, and a fragment at $m/z$ 500 is detected due to the presence of a terminal amino group at position 4′.

An LC-MS/MS method was developed for the analysis of bulaquine (BQ) 3-[1-[4-[(6-methoxy-8-quinolinylamino] pentylamino] ethylidene]dihydro-2 (3H)-furanone (Fig. 5) and its metabolite primaquine in monkey plasma. Protonated species at $m/z$ 370 and 260 were detected for bulaquine and primaquine, respectively. MS/MS conditions were optimized generating product ions through fragmentation of the molecular ions. Based on the fragmentation spectra obtained from $[M+H]^+$ under the established analytical conditions, a fragmentation pattern was presented for these compounds. This type of study is important to establish analytical conditions for the quantification of drugs and their metabolites in biological fluids. (Nitin et al., 2003)

An ESI-Ion trap mass spectrometer was used to perform MS^n analyses, in the study of imidazolidin-4-one peptidomimetic derivatives of primaquine. (Vale et al., 2008a)

4.2. Pharmacokinetic studies

Pharmacokinetic (PK) studies provide a mathematical basis to assess the time course of drug in the body. It enables to quantify absorption, distribution, metabolism and excretion of the drug and their metabolites. The primary requirement to undertake a PK study, is to have an
analytical method which is reliable, reproducible, sensitive, selective, and if possible, compatible with high-throughput pharmacokinetic approaches.

Drug efficacy requires adequate drug concentration at the site of action. Monitoring drugs and their metabolites in biological samples (in vivo studies), is fundamental in order to control the intake of the antimalarials by the infected populations. Mass spectrometry has been successfully used for this purposes, coupled with liquid chromatography, and analytical methods
have been optimized and validated for quantification of different drugs and metabolites in biological fluids.

A LC-MS/MS method was used to study 14 antimalarial drugs, which are the components of the current first-line combination treatments for malaria (artemether, artesunate, dihydroartemisinin, amodiaquine, N-desethyl-amodiaquine, lumefantrine, desbutyl-lumefantrine, piperaquine, pyronaridine, mefloquine, chloroquine, quinine, pyrimethamine and sulfadoxine). The best conditions for mass spectrometry analysis were optimized (Hodel et al., 2009) and the method developed was implemented, and used to analyse samples from an in vivo study with 125 Southeast Asian patients from two regions of Cambodia: one region with a high level of antimalarial drug resistance and another region with moderate levels of drug resistance (Hodel et al., 2010). The 14 antimalarial drugs were measured in plasma samples from the patients, and results showed that for half of them, an antimalarial drug was detected namely mefloquine, piperaquine, chloroquine or quinine. However all patients reported either not having taken any antimalarial before or not knowing to have taken. These results are important, as they show that it is urgent to ensure appropriate use of antimalarials among the populations.

Tafenoquine (8-[(4-amino-1-methylbutyl)amino]-2,6-dimethoxy-4-methyl-5-(3-trifluoromethyl-phenoxy) quinoline succinate) (Fig. 6) was measured in human plasma from patients and
healthy volunteers during a clinical efficacy trial, using a LC-MS/MS equipment. (Doyle et al., 2002)

**Figure 6. Tafenoquine**

For NPC1161 (Fig. 7), an 8-aminoquinoline analog (8-[(4-amino-1-methylbutyl)amino]-5-[3,4-dichloro-phenoxy]-4-methyl-quinoline) and their metabolites, a LC-MS method using an electro-spray ionization source and a TOF analyzer, was implemented.

**Figure 7. NPC1161, an 8-aminoquinoline analog**

Using mass spectrometry other antimalarial molecules as α-/β-diastereomers of arteether (AE), sulphadoxine (SDX) and pyrimethamine (PYR) (Sabarinath et al., 2006) and three novel trioxane antimalarials (Fig. 8) (Singh et al., 2008) were determined in rat plasma. A N-alkylamidine compound, M64, and its corresponding bioprecursors were measured in human and rat plasma (Margout et al., 2011) (Fig. 9). Artemisinin class compounds and its active in-vivo metabolites were analysed in monkey plasma (Singh et al., 2009).

**Figure 8. Chemical structures of three novel trioxane antimalarials**
A method was also developed and validated according to FDA guidelines for simultaneous
determination of two mono-thiazolium compounds in plasma, whole blood and red blood
cells from human and rat (Taudon et al., 2008). More recently a rapid (3 min analysis) and
sensitive UPLC-MS/MS method using a triple quadrupole tandem mass spectrometer in
positive ESI mode, has been implemented for the analysis of ARB-89 (7α-hydroxy artemisinin
carbamate) (Fig. 10) in rat serum, for pharmacokinetics studies, (Pabbisetty et al., 2012).

Mass spectrometry has proven to be particularly useful in identifying complex metabolites as
those arising from phase I (e.g. those involving cytochrome P450 monooxygenases) and phase
II (e.g. conjugation with glucuronic acid, sulfonates, glutathione or amino acids) reactions. In
a study published by Liu et al., 2011, metabolites of artemisinin, also known as Qing-hao-su
(QHS), and its active derivative dihydroartemisinin (DHA) were identified in \textit{in vitro} and \textit{in vivo}
biological samples using a LTQ-Orbitrap mass spectrometer in tandem with H/D ex-
change. The authors were able to show that artemisinin drugs mainly undergo hydroxylation
and loss of oxygen in the phase I metabolic pathway and can form glucuronides in the phase
II processes, as shown in Fig. 11. Based on MS data it was proposed a metabolic pathway for
these metabolites.

Piperaquine was synthesized for the first time about 50 years ago, but seems to be a suitable
partner drug in artemisinin-based combination treatments. In a paper published by Tarning
et al., 2006, the main metabolites of piperaquine were characterized in a 16-h human urine,
after a single p.o. administration of a fixed combination of dihydroartemisinin-piperaquine,
with a fatty meal. A LC method in tandem with a QTRAP system was used to analyse
piperaquine and its metabolites and a FT-ICR/MS equipment was used to obtain mass spectra

\begin{figure}
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\includegraphics[width=\textwidth]{figure9.png}
\caption{Chemical structures of compound, M64, and its corresponding bioprecursors}
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\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure10.png}
\caption{ARB-89 (7α-hydroxy artemisinin carbamate)}
\end{figure}
of the five metabolites of piperaquine in urine samples. Two of the metabolites (a carboxylic and a mono N-oxidated piperaquine metabolite) were considered as the most relevant as they were detected in the serum/plasma samples collected up to 93 days, and also in urine 123 days after administration of the drug. Other two monohydroxylated metabolites and a di-N-oxidized metabolite were also detected in urine samples.

4.3. Detection of impurities

A major impurity (8-(4-amino-4-methylbutylamino)-6 methoxyquinoline) associated with primaquine drug samples (Fig.12), obtained from European Pharmacopoeia and other commercial sources, was detected by gas chromatography-electron impact-mass spectrometry (GC–EI-MS) (Dongre et al., 2005).

4.4. Chemical stability studies

Mass spectrometry can also be conducted in order to contribute to study the properties of compounds, through knowledge of their stability and fragmentation mechanisms, under the gas-phase conditions of a mass spectrometer. This type of studies can have, in the future, important implications in drug analysis and development.
Studies on primaquine derived imidazolidin-4-ones using an ESI-ion trap mass spectrometer have allowed to find a correlation between the stability of the ions in the nozzle-skimmer region during the CID (Collision Induced Dissociation) analyses and reactivity in both isotonic buffer and human plasma (Vale et al., 2008). The same authors (Vale et al., 2008a) studied imidazolidin-4-one peptidomimetic derivatives of primaquine, PQAAPro and PQProAA (Fig. 13) and they also found the parallelism between compound reactivity to hydrolysis and stability during CID analysis. Using CID and MS/MS experiments to study the peptidomimetic imidazolidin-4-one derivatives of primaquine, it was possible to conclude that CID spectra reflected the reactivity of compounds under physiological conditions, and the relative abundances of MS/MS generated fragments were correlated with the Charton steric parameters associated to amino acid side chain in the molecule (Vale et al., 2009).

From the results obtained, ESI-MS proved to be a reliable tool for stability prediction of compounds towards hydrolysis at physiological pH and temperature although the mechanisms in water and in the gas-phase are not comparable. This type of studies were, for the first time, approached by these authors.
4.5. Detection of counterfeit drugs

The quality of commercially available drugs varies among countries. The WHO/International Medical Products Anti-Counterfeiting Taskforce estimates that in some less developed countries, the counterfeit drugs are up to 50% of the total drugs supplied to the populations (Hall et al., 2006). Due to the lack of regulations and poor quality control practices, the amount of the active ingredient may be incorrect, as a result of chemical degradation that occurs due to poor storage conditions, especially in warm and humid tropical environments. In some cases, expired drugs are repackaged with new expiration dates and put in the market. Also some drugs can be contaminated or replaced by other substances and people consume sawdust, paint and other toxic or inert substances (Kaur et al., 2009).

The report published by WHO in 2011, about the quality of antimalarial drugs in 6 countries of sub-Saharan Africa (Cameroon, Ethiopia, Ghana, Kenya, Nigeria and the United Republic of Tanzania) resumes the results obtained from the analysis of 935 samples and gives an idea about the quality and counterfeits that more often occur.

As an incorrect intake can result in a low bioavailability of the drug in the individual, leading to drug-resistance strains, provoking a therapeutic failure, reliable methods of analysis must be available to determine the quality of antimalarials commercialized, and LC coupled with mass spectrometry can be used. However this technique is expensive, requires training, technological support, and sample preparation. Actually, DART (Fernández et al., 2006) and DESI (Hais et al., 2007) methodologies are often used, as they produce results rapidly, because they do not require sample preparation. Results obtained from a DESI MS method were used to validate Fourier-transform infrared imaging for characterization of counterfeit antimalarial pharmaceutical in tablets (Ricci et al., 2007). A DESI MS method was also applied for the quantitative screening of counterfeit antimalarial tablets containing artesunate (Nyadong et al., 2008), and more recently DESI and DART methods were used to validate results from the application of FT-Raman spectroscopy for \textit{in situ} screening for potentially counterfeit artesunate antimalarial tablets (Ricci et al., 2008).

4.6. Rapid diagnosis of malaria

The rapid diagnosis of malaria infection can also be performed by mass spectrometry. Hemozoin, the malaria pigment, can be detected by laser desorption mass spectrometry (LDMS) in human blood. (Scholl et al., 2004) Detection of malaria in 45 asymptomatic pregnant Zambian women was performed by this technique. Detection of \textit{Plasmodium falciparum} malaria during pregnancy is complicated by sequestration of parasites in the placenta reducing peripheral blood microscopic detection. LDMS was able to detect <10 parasites/μL cultured in human blood and provided a more rapid mean of screening for infection than the technique used currently for this purpose, light microscopy (Nyunt et al., 2005).

5. Conclusions

The information obtained from the use of mass spectrometry in the study of antimalarials agents is important, in order to understand all the mechanisms of the illness, malaria, and the
way the different drugs interact in the human organism. Due to its characteristics (sensitivity, speed, possibility to be automated, possibility to combine with separation techniques) and diversity of equipments available, mass spectrometry can be used in the structural identification of new molecules, in the study of many phases of drug discovery, for assessment of compound stability, pharmacokinetic studies monitoring the concentrations of antimalarials and metabolites in biological matrices, the studies of cell permeability and plasma protein binding, and finally, in the quality control of commercial drugs.

The use of mass spectrometry to predict stability of compounds in physiological conditions may become an important tool.

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