Recent Advances in Real-Time Mass Spectrometry Detection of Bacteria

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Abstract
The analysis of bio-aerosols poses a technology challenge, particularly when sampling and analysis are done in situ. Mass spectrometry laboratory technology has been modified to achieve quick bacteria typing of aerosols in the field. Initially, aerosol material was collected and subjected off-line to minimum sample treatment and mass spectrometry analysis. More recently, sampling and analysis were combined in a single process for the real-time analysis of bio-aerosols in the field. This chapter discusses the development of technology for the mass spectrometry of bio-aerosols, with a focus on bacteria aerosols. Merits and drawbacks of the various technologies and their typing signatures are discussed. The chapter concludes with a brief view of future developments in bio-aerosol mass spectrometry.

1. Introduction

1.1. General
Real-time detection of biological material with absolute identity determination is the stuff of science fiction. In this case, science fiction actually represents the “market pull” that challenges a “technology push.” As concerns the real-time analysis of biological aerosols, the challenge was taken up in the mid 1980s, when some of the technology seemed sufficiently mature for integration in a universally applicable real-time bio-aerosol mass spectrometer. Two decades later, direct bio-aerosol mass spectrometry is still in the early stages of technology development, with research going on in a few select institutes. In the same two decades, the “market pull” of science fiction was fortified by an increasing awareness of biological threat, propelled by possibly emerging pandemics (e.g., Perdue and Swayne 2005; Glass and Becker 2006), and by the fear of bioterrorist attacks (e.g., Hamburg 2002; Tegnell et al. 2006). By now, mass spectrometry (MS) has proven capable of producing complicated spectral signatures from biological materials within seconds. This positions MS as a core technology for meeting the challenge of bio-aerosol detection with adequate differentiation of detected agents. This chapter gives an overview of several lines of MS technology development and of the current state-of-the-art in real-time bio-aerosol MS. After the previous two decades of early technology development, the next five years will see the evolution of mature MS-based bio-aerosol detection.

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1.2. Scope

This chapter covers the MS of aerosols, in a broad sense, with a focus on the mass spectrometry of bio-aerosols, particularly of bacteria. Viruses and other agents are not considered, mainly because there are hardly any published reports on bio-aerosol MS of such agents. This section of the introduction deals with two lines of approach: MS of bacteria and MS of aerosols, in respective subsections. The next section discusses the current state of technology of bio-aerosol MS, after a brief introduction that ties MS of bacteria to MS of aerosols. The chapter concludes with a section on the outlook, focusing on the development of fieldable bio-aerosol mass spectrometers.

1.3. MS in the Whole Cell Analysis of Bacteria

Nowadays, several forms of MS are widely employed in the typing of bacteria. The information gained by MS can range from the profiling of intact cells to the detail of proteomics or nucleotide sequencing. However, chemical analysis by MS is an emerging method with respect to established microbiological typing methods, like serotyping, microscopy, and phage typing. Where identity of organisms is discussed between analytical chemists and microbiologists, misunderstanding often arises from a difference in perception of “identity” between the two disciplines. This issue is briefly addressed in the first subsection, to clarify the point that the many approaches to the identification of bacteria correspond to various levels of certainty. The second subsection gives a brief, non-exhaustive overview of applications of MS to the analysis of whole cell material from bacteria, with emphasis on aspects of potential screening and detection technology.

1.3.1. The Definition of ‘Identity’ of Bacteria

The understanding of “identity” in microbiology differs considerably from that of chemistry. In chemistry, identity is assigned on the basis of comparison of molecular structure, for example by spectroscopic methods. Although we like to think that live material is also subject to the rigidity of molecular structure, the adaptive and evolutionary qualities of life imply a certain fuzziness in chemical composition. Historically, classical microbiology and bacteriology set out to assign microorganism identity by descriptive comparison of a selected set of observable characteristics. These characteristics would be morphological, for example colony shape in culturing or cell shape in microscopy. Later methods of typing employed indirectly observable characteristics, for example particular enzymatic activity, chemical resistance or affinity, or immunological activity. With the advent of chemical methods of analysis, typing would also address parts of the chemical composition of microorganisms, like fatty acid distribution or excreted metabolites. The molecular biology revolution, which started around 1980, has provided typing methods based on hereditary properties as these are addressed through the analysis of DNA. It is envisaged that molecular biology may provide a Grand Unification Theory for typing of microorganisms. That will also resolve the practical problem of the evolution of “standards” because the abstract information can be documented on paper rather than in a collection of organisms. However, where the link between DNA sequence and biological function is still largely elusive, molecular biology cannot yet accomplish unification by the incorporation of existing typing schemes.

Without that Grand Unification scheme accomplished or even partly available, every bacterial typing method occupies its own dimension (see, for example, Spiegelman, Whissell, and Greer 2005). Studies employing multiple typing methods provide a link between the separate dimensions, often on an empirical basis. For example, if a bacteriophage susceptibility profile is obtained from a specifically serotyped Salmonella isolate, a single mapping term...
is established between bacteriophage typing and serotyping. If mapping from a given typing method to other typing methods is not available, that typing method remains subjective. For practical purposes, for example in epidemiological comparison of bacterial strains from an outbreak, that subjectivity may be sufficient. When the identification problem is limited to a “threat list,” subjectivity will be insufficient. Even when the typing profiles of all organisms on the “threat list” are acquired, it remains to be proven that no organisms on the list give the profile of one of the threat agents. For typing in any environment, a link to objective information, generally a DNA sequence, is eventually required to overcome the typing method dimension problem.

As MS methods for the typing of bacteria are concerned, the distance to the objective information of genomic DNA can be qualitatively evaluated. Fig. 36.1 gives the relative information distance of several characteristics of bacteria as they are accessible by chemical analysis.

1.3.2. Mass Spectrometry of Bacteria

Mass spectrometry is currently widely employed in the analysis of bacteria or their constituent compounds (as reviewed, see Van Baar 2000; Fenselau and Demirev 2001; Lay 2001; Fox 2006). However, many of the laboratory methods are not suitable for the detection of aerosolized bacteria, because sample treatment is too elaborate. Therefore, analysis of whole cells with little or no sample treatment is the only option, with the process applied to bulk material or to single cells in single particles. This subsection gives an overview of whole cell MS of bacteria, with emphasis on the signature obtained.

1.3.2.1. Pyrolysis MS

Early studies typically employed the mixture analysis capabilities of the then-newly emerging GC-MS technology, in pyGC-MS (Simmonds, Shulman, and Stembridge 1969; Simmonds 1970). A transition to direct inlet pyMS was quickly made for detection purposes because GC separation is relatively time consuming (Meuzelaar and Kistemaker 1973). Also, the use of field ionization (FI) (Schulten et al. 1973), chemical ionization (CI; Van der Greef, Tas, and Ten Noever de Brauw 1988), and metastable atom bombardment (MAB; Wilkes et al. 2005) with pyMS was explored as an alternative to common electron ionization (EI).
Other studies addressed spectral comparison, for example through multivariate methods (Shute et al. 1984), artificial neural networks (Freeman, Sisson, and Ward 1995), or genetic algorithms (Taylor et al. 1998). PyMS was successfully applied in the differentiation of pathogens within specific families of bacteria, for example *Mycobacteria* (Wieten et al. 198) and *Salmonellae* (Van der Greef, Tas, and Ten Noever de Brauw 1988; Wilkes et al. 2005). Even though specific compounds were identified as pattern markers (e.g., DeLuca, Sarver, and Voorhees 1992), identification has always remained confined to a “fingerprinting” approach requiring the availability of a library or of a basis set of qualified spectra. Therefore, pyMS of bacteria is still typically used for quick screening, for example of clinical samples (Kyne et al. 1998; McCracken et al. 2000) or mail (Wilkes et al. 2006).

For many purposes, comparative matching of spectra or profiles is unsatisfactory because it does not allow a translation to other and independent knowledge available on the biological material at hand. For bacteria, this problem was approached by the application of MS/MS or by a minimum sample pretreatment. An example characteristic accessible by MS/MS is dipicolinic acid (DPA). This compound makes up 5-15% of a *Bacillus* spore, and as such it is representative for the possible presence of *Bacillus anthracis*. Specific pyMS based methods for detecting *Bacillus* through DPA were developed (Beverly et al. 1996; Goodacre et al. 2000; Tripathi, Maswadeh, and Snyder 2001). Tandem mass spectrometry was employed to show that a more general characteristic of bacteria, their fatty acid profile, is accessible by pyMS/MS without prior chromatographic separation (DeLuca et al. 1990). A fast *in situ* chemical methylation reaction, to form fatty acid methyl esters (FAMEs), is included to allow better volatilization (Basile et al. 1998; Barshick, Wolf, and Vass 1999; Tripathi, Maswadeh, and Snyder 2001; Poerschmann et al. 2005). Although FAME profiling does not provide a complete typing scheme for bacteria, correlation with laboratory profiles can be employed in detection applications. Thus, the targeting of specific components by pyMS through MS/MS or fast *in situ* sample treatment obviates the need for elaborate full spectrum matching in detection.

### 1.3.2.2. LDI MS

LDI or LAMMA have not found much application in the typing of bacteria. The elemental analysis capability of LAMMA was employed to measure the Na⁺/K⁺ ratio in individual cells in order to establish the live or death status of *Mycobacteria* in response to external factors (Seydel et al. 1982; Seydel and Lindner 1988). Böhm et al. (1985) demonstrated laser ionisation of single bacterial cells from three *Bacillus* species. They concluded that high laser power densities (>1000 Wcm⁻²) were required to obtain positive or negative ions in pyMS-like spectra (see Fig. 36.2).

A discriminant analysis showed that typing by classification was poor, 70 to 75%, even with this small *Bacillus* training set (Böhm et al. 1985). One later study, which compared several desorption/ionization methods in the analysis of cells, showed that the type of laser ionization employed in LDI and LAMMA gave signals of intact polar lipids from lysed *E. coli* (Heller et al. 1987). From these pioneering studies it is clear that LDI and LAMMA do not provide a sound basis for bacteria typing, because typing information is at best highly convoluted and of poorer quality than the information obtained from pyMS. In addition, the first reports on application of MALDI MS to bacteria, in the mid 1990s, left LDI and LAMMA obsolete as far as bacteria typing was concerned.

### 1.3.2.3. MALDI MS

Three pioneering studies on MALDI MS of vegetative whole bacteria were published independently in 1996 (Holland et al. 1996; Claydon et al. 1996; Krishnamurthy and Ross
Figure 36.2. Typical LAMMA mass spectra of *Bacillus cereus* from pioloform foil with a copper grid, at a laser power density of 85% (top) and 35% (bottom) of the full Nd:YAG laser power (reprinted from Böhm et al. (1985), with permission of Elsevier).
Three different sets of bacteria were employed in the three studies to arrive at a similar conclusion: Specific biomarker signals were repeatedly observed in the mass spectral profiles. Moreover, these signals allowed typing of the bacteria, at genus, species and strain level, by comparison to earlier recorded spectra or to newly acquired spectra from freshly cultured strains (see Fig. 36.3). Although these three studies used different mass ranges and different methods for sample application to the MALDI target, the work established a starting point for later studies.

The robustness of whole cell MALDI MS of bacteria was tested in many studies, which all show that sample preparation and details of the ionization process are important parameters for experimental control (e.g., Wang et al. 1998; Evason, Claydon, and Gordon 2001; Saenz et al. 2001; Williams et al. 2003). Bacterial growth conditions are beyond experimental control, at least in a real detection environment, but their effect on the variability of spectra has triggered further study. For example, it was observed that number and intensity of peaks varied considerably among spectra when *E. coli* K12 bacteria were sampled at selected stages between 6 to 84 h after inoculation of a culture (Arnold et al. 1999). However, the general trend is that protein fingerprinting in whole cell MALDI provides adequate identification despite the innate variability (e.g., Valentine et al. 2005; Wunschel et al. 2005a). Several studies with a wide scope have been reported, for example an interlaboratory study (Wunschel et al. 2005b) and some large scale typing studies for food pathogens (Mandrell et al. 2005), clinical isolates (Nilsson 1999; Marvin, Roberts, and Fay 2003; Keys et al. 2004), and in biotechnology (Jones et al. 2005).

Data analysis is an important and yet unsettled issue in whole cell MALDI MS. Most of the data analysis research focuses on the processing of spectra as “fingerprints.” Because similar fingerprint type processing is also used in pyMS, procedures can be adapted for the purpose. However, plain spectrum fingerprint matching discards the more hidden information from the bacterial genome. The observed high-mass signals, over 1000 Da, are generally taken to represent peptides and proteins of which the mass is determined by the amino acid sequence. Therefore, these signals are considered “biomarkers” of the investigated bacteria. The biomarker link to genomes provides a more objective kind of information (Demirev et al. 1999), from a sequence database that is essentially independent of any analytical method. The notion of biomarkers implies that there is a potential for the kind of spectrum prediction that obviates the need for library accumulation. At present, fingerprinting is the established way for bacteria
typing by MALDI MS, and the link to genome information is still under investigation. Both approaches are briefly discussed below.

Fingerprints may be accumulated as an internally consistent set, while the actual spectra are machine-dependent and sample dependent. Raw spectra are rarely used as fingerprints, and several condensation methods have been reported, for example cross-correlation (Arnold and Reilly 1998) and extraction of numerical fingerprints (Jarman et al. 1999; Jarman et al. 2000). Actual spectrum comparison is then achieved by methods also employed in matching of pyMS spectra, for example in the neural network approach demonstrated with SELDI (Schmid et al. 2005). The accumulation of a well-defined set of library spectra, possibly in a condensed form, is an elaborate job. However, it is generally assumed that translation of fingerprint data to other instruments and conditions is possible. This assumption was studied in an objective way by consideration of the protein distribution against the significance of an identification, for a limited set of organisms (Pineda et al. 2000). It was concluded that the cluttered and incomplete nature of the spectral data, as compared to bacterial proteomes, complicates or even compromises truly robust identification. At a practical level, the assumption of cross-instrument translation is supported by a single published interlaboratory study, which showed that tight control of sample treatment and of instrument conditions allows translation of MALDI MS fingerprints between instruments (Wunschel et al. 2005b). The assumption is further corroborated by the development of two commercially available MALDI MS platforms with libraries for bacteria typing: from Waters (employed, e.g., by Keys et al. 2004) and from Bruker (employed, e.g., by Maier et al. 2006). Although theoretical considerations and practical experience have not yet fully resolved the issue of method robustness, bacterial fingerprinting has become fairly well established as a method for bacteria typing in a laboratory setting and with a certain level of prior knowledge about target bacteria, for example in the screening for dairy pathogens.

Although the word biomarker appears in the pioneering papers on whole cell MALDI MS of bacteria, the link of signals to actually identified bacterial proteins was fairly speculative or even absent. Later studies provided more substantial evidence, for example for the identity of certain proteins from Bacillus spores (Hathout et al. 1999) or independent proof of protein identity for specific MALDI MS signals (e.g., Hathout et al. 2003; Dickinson et al. 2004). On the basis of this sporadic approach it was proposed by the group of Fenselau that it would be most likely that basic proteins, like many of the ribosomal proteins, are responsible for biomarker signals (Pineda et al. 2003). The proposal was corroborated by experiments that showed it is tenable as a working hypothesis, provided that post-translational modification of the proteins is accounted for (Demirev et al. 2001). Although this working hypothesis does not have any absolute predictive value, the basic protein approach is the most generalized proposal for biomarker attribution to date.

### 1.3.2.4. Comparison of MS Methods for Whole Cell Bacteria Typing

PyMS, LDI MS and MALDI MS have proven useful in rapid typing of bacteria in the laboratory, for which purpose they are operated with bulk samples and in a batchwise fashion. No direct experimental comparison of these methods is available. In addition, differences in ionization, in the bacterial compounds addressed, and in possible sample treatment hamper a theoretical comparison, for example of the sensitivity. For a qualitative consideration, a comparison can be made of the capability for generic bacteria typing to the attainable degree of resolution in typing (to the increasingly more detailed levels of bacteria/non-bacteria, bacteria genus, species, strain, and isolate).

PyMS, as it is applied in rapid analysis, will give information about the presence of a Bacillus genus marker (DPA), or it will produce a FAME profile. As a Bacillus detector, pyMS is not generic, and it has no further degree of typing resolution. From laboratory studies of FAME profiling without pyMS but by plain GC or GC-MS (e.g., Abel, DeSchmertzing, and
Peterson 1963; Moss 1990), it is clear that typing resolution may be highly specific within certain families of bacteria and fairly unspecific in other families. In addition, FAME profiles are known to change with changing environmental conditions, such as bacterial nutrition (see, e.g., Stoakes et al. 1991). As a FAME profiler, pyMS is a generic method, because such a profile can be generated from any bacteria. Overall, pyMS does cover some of the needs for rapid typing in the laboratory, while the method seems to have reached its full potential.

LDI MS and LAMMA, as they can be applied in rapid analysis, will not give very distinctive information. The near-atomization conditions of the direct laser ionization destroy the structure of biological compounds to an extent that highly specific molecular information is not observed. No recent reports have appeared on LDI MS or LAMMA investigations of bacteria in the laboratory. For laboratory typing of bacteria, LDI MS and LAMMA have been surpassed by MALDI MS at the present state of technology.

MALDI MS, as it is applied in rapid whole-cell analysis, will produce a biomarker profile in a single mass spectrum. Because this procedure can be done with any bacteria, the method is generic. Opinions on the specificity and usefulness of such a profile have come to some consensus with an increasing number of studies becoming available. The general conviction (see, e.g., Lay 2000) is that certain true biomarkers will always be present, whereas other biomarker signals will show major variations with environmental conditions such as culturing or growth stage. The principal advantage that biological molecules stay intact in MALDI, to retain the information content, is hardly employed, because most studies use plain profile matching. Overall, MALDI MS is well suitable to rapid typing of bacteria in the laboratory, while the method requires further exploration to assess its full potential.

1.4. Aerosol MS

MS analysis of aerosols can be accomplished in many ways. Some of the typical laboratory MS methods have been adapted to allow analysis of deposited aerosols, with sampling and chemical analysis offline. In addition, dedicated aerosol MS analysis technology was developed, to allow direct analysis of single particles from atmospheric aerosols without sample deposition. Although the deposition and direct analysis type methods share some of the technology, they are discussed separately.

1.4.1. MS of Deposited Aerosols

Bio-aerosols are often made amenable to mass spectrometry by particle collection. Although particle collection will allow further preparation for any desired MS analysis, subsequent sample treatment is generally minimized to allow quick analysis. Currently, one of three modes of MS analysis is typically employed: pyrolysis with EI or CI, laser desorption/ionization (LDI), and matrix-assisted LDI (MALDI). Because these modes of analysis produce distinct information from biological material, they are briefly discussed.

1.4.1.1. Pyrolysis MS

Pyrolysis generates volatile compounds from biological material, both by evaporation of any volatile compounds present and by the formation of new volatile compounds in elimination reactions under the influence of heat. The pyrolysis process can be conducted in a classical MS source to allow subsequent ionization. Many mass spectra can be acquired during a pyrolysis cycle to give a time-resolved profile. The spectra, the profile, or both can be employed as a characteristic for that particular biological material.

Pyrolysis MS (pyMS) initially evolved as a method for the chemical analysis of bacteria after it had been shown in the 1960s that pyrolysis gas chromatography (pyGC) provided
“fingerprinting” of bacteria (see, e.g., Oyama 1963; Reiner 1965; Stern, Kotula, and Pierson 1979). Therefore, pyMS of bio-aerosols is discussed in more detail in the next section.

1.4.1.2. LDI MS

Laser desorption/ionisation (LDI) was developed in the late 1970s and early 1980s (e.g., Hillenkamp et al. 1975; Stoll and Röllgen 1979; Cotter 1980), when suitable laser technology became available. LDI was typically used for samples applied to a surface, and to direct surface analysis. In LDI, the ionizing laser energy is dissipated by the sample material and the corresponding energy density is generally so high that covalent chemical bonds easily dissociate. This laser ablation process burns away minute amounts of sample material. Therefore, a laser desorption spectrum will generally show signals of native ions such as Na$^+$ and NO$_3^-$, and of atomic cluster ions such as C$_3^+$ and C$_3$H$_-$, formed from the organic or biological compounds present. A typical example of such a spectrum is shown in Fig. 36.4. The high energy density in LDI makes the method less suitable for the ionization of relatively thermolabile organic and biochemical compounds. Nevertheless, occasional reports of successful LDI analysis of such compounds have appeared (see, for example, Balasanmugam et al. 1986; Posthumus et al. 1978). In many cases, the ionization of such thermolabile compounds involved preformed cationized and anionized species (Cotter 1981; Balasanmugam et al. 1981; Zakett et al. 1981).

The utility of LDI was greatly enhanced by combining the ionization with microscopic accuracy control of the ablation position to within 1 µm resolution (Hillenkamp et al. 1975; Wechsung et al. 1978). That form of LDI, generally known as Laser Ablation Microprobe Mass Analysis (LAMMA), was typically used for the analysis of samples on a surface. Initially, the capability for analysis of deposited aerosol particles was oddly reported among biomedical applications (Kaufman, Hillenkamp, and Wechsung 1979), but LAMMA was then quickly adopted by aerosol investigators (Wieser, Wurster, and Seiler 1988). The initial studies concerned technology development, for example for measuring the elemental composition of micron size particles (e.g., Bruynseels and Van Grieken 1984). This technology was then applied in atmospheric aerosol research (e.g., Bruynseels et al. 1988; Dierck et al. 1992; Hara et al. 1996). These aerosol studies generally encompassed inorganic analysis, where the particle
size range investigated was set by the aerosol fraction trapped by the offline sampler. In these studies, LAMMA MS is typically used for the analysis of the low Z elements, such as C, N, and S. Other methods, like electron probe X-ray microanalysis (EPXMA), turned out to be more suitable for the higher Z-elements, such as heavy metals (e.g., Van Malderen, Hoornaert, and Van Grieken 1996). Although only a single LAMMA study was reported in recent years, the available aerosol sampling technology and offline LAMMA analysis in a remote laboratory provided many new insights in environmental aerosol chemistry.

1.4.1.3. MALDI MS

MALDI MS evolved from LDI in the late 1980s (Tanaka et al. 1987; Karas and Hillenkamp 1988), when it turned out that the application of a specific “matrix” compound with a sample produced high mass ions upon UV laser irradiation. The matrix is thought to dissipate most of the incident UV energy. The matrix is also supposed to be a proton donor in positive ionization, and a proton acceptor in negative ionization. Although several studies were devoted to elucidation of the details of the MALDI ionization process (see, e.g., Knochenmuss and Zenobi 2003; Karas and Krüger 2003), there is no consensus theory or model. As a consequence, much of the chemistry involved in MALDI is still empirical.

The persistence of the empirical experimental component has not kept MALDI from gaining enormous acclaim as a method for MS of biological compounds. Empirical studies in MALDI MS covered issues like matrix crystallization (e.g., Westman et al. 1995; Dai, Whittal, and Li 1996; Luxembourg et al. 2003) and sample desalination (e.g., Kussman et al. 1997), to set the standard for a plethora of applied studies of large molecules in polymer science (e.g., Nielen 1999; Macha and Limbach 2002), microbiology (viruses, e.g., Lewis et al. 1998; Fenselau and Demirev, 2001; for bacteria, see above), and direct tissue analysis and imaging (e.g., Stoeckli, Farmer, and Caprioli 1999; Schwartz, Reyzer, and Caprioli 2003; Altelaar et al. 2005). MALDI MS also found application in aerosol analysis, as will be discussed later on.

For the purpose of this overview, surface enhanced laser desorption/ionization (SELDI) should also be included. SELDI was developed as a combination of MALDI with affinity capture chip technology on the target (Hutchens and Yip 1993; Caputo, Moharram, and Martin 2003). SELDI is the sophisticated, often commercialized variant of earlier academic experiments with on-target sample preparation (e.g., Brockman and Orlando 1995; Liang et al. 1998; Bundy and Fenselau 1999) for MALDI. The SELDI chip that serves as the laser target is coated with specific affinity molecules to allow on-target cleanup before matrix application and actual MS. Mass spectra are then obtained from any material retained by the affinity surface, for a rapid characterization of target material captured from the original sample. The chip technology makes SELDI particularly useful for quick screening of large amounts of samples, for example in clinical diagnosis of cancer by biomarkers (recently reviewed in Engwegen et al. 2006) and for bacteria typing (e.g. Schmid et al. 2005; Al Dahouk et al. 2006).

1.4.2. Direct MS of aerosols

In the 1980s LDI MS and LAMMA were proven to be of use in atmospheric (bio-)aerosol research. However, the required particle collection provided a cross-section picture of a certain particle size class, but any particle concentration information got lost in the process. In a separate development, Sinha et al. constructed an aerosol mass spectrometer for the analysis of individual particles. A beam of particles was continuously fed into the source of a quadrupole (Q) mass spectrometer, with in-source pyrolysis by a heated filament and electron ionization (Sinha et al. 1985). A key problem is that Q type mass analyzers require a constant ion input for a time span in the order of one second during a full mass range scan, whereas single particle
ionization produces only a short burst of ions during a period in the order of a millisecond. This problem was resolved by using the Q mass spectrometer as a band pass filter at a width of a few mass units. By stepping through multiple bands the 40–300 Da mass range was covered. This yielded mass spectra from a bacteria aerosol (Fig. 36.5), where a single spectrum would have required ionization of over $10^5$ bacteria containing particles. This kind of sensitivity and mass range are not practically useful, but the work provided a most important proof of principle.

The concepts of the use of laser ionization (Sinha 1984) and of the use of time-of-flight (TOF) instead of Q mass spectrometers (Marijnissen, Scarlett, and Verheijen 1988) for single particle analysis were clear by the mid 1980s. However, the actual hardware development took until the early 1990s, when suitable lasers and mass spectrometers had finally become available. The technology was discussed in several reviews (Suess and Prather 1999; Noble and Prather 2000; Murphy 2006) and a summary description is given in the next section before the current state of the technology is addressed.

### 2. Current State of the Technology

#### 2.1. Considerations on Aerosol MS of Bacteria

From the technology for aerosol MS and the application of MS to non-aerosolized bacteria it follows that the routine use of aerosol MS for the detection of bacteria is within reach of
technology. It is also clear that the development and performance of such bio-aerosol MS technology is determined by several issues:

- aerosol deposition versus direct aerosol analysis;
- the possibility of applying any degree of sample treatment in the analysis process, either to single particles in the aerosol phase or after aerosol deposition;
- access to genome or proteome linked information versus accumulation of fingerprint type information.

At present, the field of bio-aerosol MS encompasses a variety of technology solutions, each of which deals with the above issues in their own way. This section discusses the available technology, as it was developed by the various research groups, with a clustering of similar technology.

2.2. Deposition and PyMS Based Technology

Pyrolysis MS based technology for the detection of bio-aerosols is currently best developed. PyMS analysis implies deposition of aerosol particles in a bulk, which process makes the particulate matter amenable to any form of elaborate sample treatment prior to the actual MS analysis (e.g., Szponar and Larsson 2001). However, because time is of the essence in bio-aerosol detection, two rapid methods with minimum sample treatment have found their way to field application: DPA confirmation and in situ derivatization with FAME profiling.

The proved presence of DPA in spores of the *Bacillus* family (see above) opened the way to a detection method for *Bacillus anthracis*, a main threat agent. Pyrolytic liberation of the acid and subsequent MS/MS in an ion-trap type instrument formed the key “biological” capability in the first version of the “Chemical and Biological Mass Spectrometer” detector (CBMS). The in situ methylation used for FAME profiling also methylates DPA. Thus, detection of dimethyl-DPA is still a key feature of the second version of the CBMS detector, the CBMS Block II (Hart et al. 2000; Lammert et al. 2002; see Fig. 36.6). Although the scope of DPA detection is limited to *Bacillus* species, it presently supports the only operational detector with a capability to differentiate to the level of an anthrax or closely related threat.

The development of a rapid process for in situ methylation, by heating with tetramethylhydroxide (TMH), formed the basis for FAME profiling in a field setting. This required some

**Figure 36.6.** Example of fieldable pyMS based equipment: the CBMS Block II instrument.

| Size          |        |
|---------------|--------|
| Height        | 91 cm  |
| Width         | 51 cm  |
| Depth         | 36 cm  |
| Weight        | 77 kg  |

| Power         |        |
|---------------|--------|
| Peak          | 1000 W |
| Continuous    | 500 W  |

**Operating conditions**

| Temperature   | −32 – +49 ºC |
|---------------|--------------|
| Humidity      | 5 – 95 %     |
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instrument and process optimization to allow sequential TMH treatment and pyrolysis in a single reaction vessel and to allow batchwise sample introduction. Several potentially transportable MS based detector systems have been under investigation for FAME profiling (e.g., Hart et al. 1999; Gardner et al. 2005), and the method has become the key mode of “biological detection” in the second version of the CBMS detector (Hart et al. 2000; Lammert et al. 2002). Some validation studies of the CBMS Block II system have appeared, to demonstrate much of its capabilities (e.g., Luo et al. 1999; Griest et al. 2001). For bacteria and fungi, the CBMS is presently the single operational biological detection system with a differentiation capability to the genus level and, sometimes, to the species level.

2.3. Deposition and MALDI MS Based Technology

The sample preparation process for on-target MALDI does not lend itself to easy automation in a batchwise process. To our knowledge, there is only a single line of development based on off-line aerosol entrapment and subsequent MALDI MS. In the mid 1990s it was demonstrated that a TOF mass spectrometer with a flight tube of a few centimeters long can have the performance of an instrument with a flight tube of over 1 meter in length (Cornish and Cotter 1992; Bryden et al. 1995). This small TOF mass spectrometer was the basis for interfacing with aerosol sampling and MALDI, as it was developed at the Johns Hopkins University Applied Physics Laboratory (JHU-APL) and described in a patent (Anderson and Carlson 1999; Anderson et al. 2003). Typically, aerosol is sampled onto a tape by an aerosol sample collector, where the sampling determines the overall duty cycle for detection. After completion of a sampling/deposition cycle, a matrix is applied and the tape is interfaced with the mass spectrometer by a sealable opening. With the sample spot on the tape in the correct position inside the mass spectrometer source, a common MALDI MS experiment can be conducted. Investigations on the performance of this system were not published in peer-reviewed journals, but aspects of system development were covered in JHU publications (e.g., Antoine et al. 2004). Although it is hardly possible to come to an evaluation of sensitivity and performance of this aerosol MALDI TOF MS system on the basis of the scant literature data, it is obvious that the technology has the full potential of MALDI MS for the identification of bacteria.

2.4. Single Particle LDI MS Technology

Following the pioneering work of Sinha (1984; Sinha et al. 1985) and a first concept of a single particle aerosol mass spectrometer (Marijnissen, Scarlett, and Verheijen 1988), first reports of actually built instruments appeared in the early 1990s (McKeown, Johnston, and Murphy 1991; Kievit et al. 1992). A typical design for a single particle MS instrument is given in Fig. 36.7 (Van Wuijckhuijse et al. 1998). Environmental aerosol is sampled into the mass spectrometer vacuum through a differentially pumped beam generator. The emerging particle beam is made to pass a low power continuous wave laser beam, where every single particle gives light scattering. The scattered light is detected and employed to fire a high power pulsed excimer UV laser, in order to ionize the particulate material. As the ionization events are located in the source of a TOF mass spectrometer, the ionization laser shot is also used to mark the start of the TOF MS process. Because this experimental set-up only allowed analysis of particles with a certain pre-set size, the demonstrator instruments served as a starting point for further exploration, pursued by several research groups.

The integration of aerodynamic sizing with triggering of the ionization laser, in a three-beam laser arrangement, made particle analysis more versatile (Weiss et al. 1993; Prather, Nordmeyer, and Salt 1994; Carson et al. 1995; Kievit et al. 1996). A double low power
laser beam passage made ionization laser triggering a size independent process, and particle beam generation now became the main determinant for the particle size range. In addition, synchronous positive and negative ion TOF MS was implemented to improve detection capability (Hinz, Kaufmann, and Spengler 1994). Redesign to a transportable instrument was investigated (Gard et al. 1997). Also, the use of Ion Trap MS, instead of TOF MS, was explored (Dale et al. 1994; Yang et al. 1995). Finally, the TOF based technology was developed into a commercial instrument for the analysis of chemical aerosols (TSI Inc., USA; www.tsi.com).

Recently, it was shown that the same technology can be applied in the analysis of single particles of a bio-aerosol composed of *Bacillus* spores or *Mycobacteria* (Steele et al. 2003; Fergenson et al. 2004; see Fig. 36.8). Ionization was accomplished with the help of a 266 nm pulsed UV laser, for which it was considered that 266 nm coincides with the UV absorption maximum of the dipicolinic acid that is abundant in such spores. In a later study it was shown that the fluence threshold and the beam profile of the ionization laser are of prime importance to the spectrum quality and abundance (Steele et al. 2005). Data analysis in the *Bacillus* study showed that \( m/z \) 74\(^+\) and \( m/z \) 173\(^-\) ions provided the main discriminant signals, where the positive ions were proposed to originate from trimethylglycine (Srivastava et al. 2005) and the negative ions from free arginine (Fergenson et al. 2004). However the robustness of these marker signals turned out to depend on the micro-organism growth conditions: Different growth conditions lowered the probability of detection of *B. atrophaeus* from 93 to 73 \%, whereas the distinction from *B. thuringiensis* was lost (Fergenson et al. 2004). Specifically for *B. atrophaeus*, the impact of growth on the particle mass spectrum was investigated in more depth (Tobias et al. 2006). It was also shown that *Mycobacterium tuberculosis* can be distinguished from *Bacillus cereus*, *B. atrophaeus*, and *Mycobacterium smegmatis*, with a single marker signal as the determinant (Tobias et al. 2005). The observed negative ion marker, \( m/z \) 421\(^-\), was tentatively attributed to a sulfolipid precursor earlier identified by independent methods as a component of the *M. tuberculosis*. Although there are just a few of these micro-organism studies, it is obvious that the focus in research on single particle LDI MS has shifted from technology exploration and development to the originally envisaged applications.

These few studies with biological material show that single particle LDI MS still holds some promise for bio-aerosol analysis. The sensitivity seems to be sufficient to give marker
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Figure 36.8. Combined axis positive and negative ion mass spectra obtained from averaging single particle LDI spectra of B. thuringiensis and B. atropheus after culturing on different growth media (Reprinted with permission from Fergenson et al. 2004. Copyright 2004 American Chemical Society).

signals from single particles. As the spectrum quality is concerned, there seems to be a strong analogy with the original pyrolysis mass spectra of bacteria, obtained in the 1970s and 1980s. Spectra typically display complicated patterns, with variable reproducibility and one or few biomarker signals. Many more microorganisms will have to be investigated to get to a more definite picture of single particle LDI capabilities.

2.5. Single Particle MALDI MS Technology

The step from single particle LDI MS to single particle MALDI MS seems logical by the historic analogy of the development of LDI and MALDI for bio-analysis. For single particle MS, this step is relatively easy when a matrix compound is mixed with the sample, prior to aerosolization. This was demonstrated with aerosols of bradykinin, gramicidin S, and myoglobin and α-cyano-4-hydroxycinnamic acid matrix (Murray et al. 1996), of leucine and of peptides (enkephalin and gramicidin S) mixed with several matrix compounds (Mansoori, Johnston, and Wexler, 1996) and with aerosols of gramicidin S with 3-nitrobenzyl alcohol as the matrix (Weiss 1997). At that time, the mass range and some other equipment limitations of typical single particle LDI mass spectrometers did not allow a full exploration of the premixed matrix
approach. In addition, premixing of the matrix had little relevance in environmental aerosol research and in bio-aerosol detection. However, in the development of bio-analytical MS for laboratory applications there is a need for online coupling of MALDI MS with liquid flows from continuous separation methods (Foret and Preisler 2002; Gelpí 2002), for example with liquid chromatography (Murray et al. 1996; Preisler, Foret, and Karger 1998; Miliotis et al. 2000) or capillary electrophoresis (Zhang and Caprioli 1996). Therefore, the development of single particle MALDI MS follows two lines of sample treatment, either with matrix premixing prior to aerosolization or with online, in-flight coating of aerosol particles.

With the focus on bioaerosol mass spectrometry, interfacing of liquid based separations with single particle aerosol MALDI MS provides some insight into the performance of potential bio-aerosol mass spectrometer equipment. With various matrix compounds and several UV wavelengths employed, it is reported that the required matrix-to-analyte ratio lies between 10:1 and 1000:1 (Beeson, Murray, and Russell 1995; Mansoori, Johnston, and Wexler, 1996; He and Murray 1999). Although the quantity in a single particle of pure analyte lies between 100 amol and 10 fmol, the analyte loss by interfacing through aerosolization implies that the overall analysis method is relatively insensitive. However, this analyte loss depends strongly on the design of the interface, and actual numbers for the analyte loss are not available. Because MALDI MS of deposited material does not ionize all of the deposited analyte, aerosol MALDI MS is still able to compete favorably. At present, chromatography interfacing to mass spectrometry is still in the research stage of development.

Analysis of environmental bio-aerosol particles by single particle MALDI MS requires in-flight coating. Particle coating has found application in existing technology (see, e.g., Agarwal and Sem 1980). In principle, matrix coating by condensation can be accomplished by subsequent particle passage through a warm saturated vapor and a cold zone (Fig. 36.9). Upon cooling, the oversaturated vapor condenses on the aerosol particles, which are the only available condensation nuclei. These coated particles are then analyzed by a single particle TOF mass spectrometer (Stowers et al. 2000; Jackson, Mishra, and Murray 2004). In the coating process, the final particle size depends on the initial size, the temperature difference between hot and cold zone, the aerosol number concentration, and the sampling rate. These parameters were employed to tune the matrix-to-analyte ratio to produce MALDI mass spectra of acceptable quality. In a proof of principle, spectra were obtained from Gramicidin S and from Bacillus subtilis var niger (known as BG), and BG signals were tentatively assigned to peptidoglycan typical for BG cells (Stowers et al. 2000). This proof of principle showed that single particle MALDI TOF MS is possible.

Figure 36.9. Schematic of an in-flight matrix coating apparatus for aerosol particles (from Van Wuijckhuijse et al. 2005b; reproduced with kind permission of Springer Science and Business Media).
The initial equipment was augmented with new ion optics to extend the mass range, and with a triggering mechanism for selective ionization of biological particles (Stowers et al. 2002; Van Wuijckhuijse et al. 2005a and 2005b). The earlier observations for BG were reproduced and additional signals were found in the 6000-7500 Da mass range. The improvements allowed registration of mass spectra in the 1 to 30 kDa range found relevant to whole cell bacteria typing in common MALDI (see above).

**Figure 36.10.** Spectra from “crushed crystallized” *E. coli* obtained on a common MALDI MS instrument (top trace; Biflex III MALDI mass spectrometer 337 nm ionization laser, average of 100 shots; Bruker Daltonics; Bremen, Germany) on two different direct MALDI ATOFMS instruments (middle: Delft University of Technology instrument with 308 nm ionization; bottom: instrument at TNO, with 337 nm ionization; both average of ∼1000 single particle spectra); vertical lines represent the mass spectrum peak list.

**Figure 36.11.** Direct Aerosol MALDI TOF mass spectra from *B. globigii* (top), *B. thuringiensis* (middle), and *B. cereus* (bottom) spores, obtained from the aerosolized spores and with spectrum averaging of ∼500 particles.
Over the last year, work has been going on to improve matrix coating, to become reproducible, and to build a prototype transportable system (Lok, 2007; Van Wuijckhuijse et al. in preparation). This will show whether single particle MALDI MS gives signatures that compare to those from whole cell MALDI MS in the laboratory. In a first step, *Escherichia coli* was prepared for a laboratory MALDI experiment by the common addition of matrix. Part of the crystallized material was collected and crushed to allow subsequent aerosolization. As it is shown in Fig. 36.10, analysis of this material with two different aerosol TOF mass spectrometers and with a common laboratory instrument yielded similar spectra. Although this single experiment requires repetition with a few more organisms, the spectrum resemblance demonstrates that spectrum matching of spectra from a single particle MALDI MS instrument with those from common whole-cell MALDI MS will support bacterial typing. The improved performance of the present system (Van Wuijckhuijse in preparation) is also demonstrated by single particle MALDI MS averaged spectra from several *Bacillus* species: *B. thuringiensis*, BG, and *B. cereus* (Fig. 36.11). These 500 particle average spectra, evaluated in their entirety and not yet by biomarker signals, clearly discriminate between the three closely related *Bacillus* species, and they also discriminate between spores and vegetative cells.

3. Conclusions and Future Perspectives

Bio-aerosol MS is still in the early stages of development, with 10 to 20 research groups world wide investigating the various technologies. In general, technology development tends toward increasing transportability, ruggedness, and ease of operation. That tendency is driven by the potential applications, which all involve *in situ* measurement for quick and reliable detection: medical, environmental, as well as civil and military safety applications. Where potential military and civil safety bio-aerosol MS is pretty much limited to the scope of threat perception, medical and environmental bio-aerosol MS may open up a completely new view of the world. We will become able to “see” microorganisms in environments where we have been blind until now. Of course, this sense of science fiction translates into some practical requirements for science and technology development. In this outlook we do not pretend to be exhaustive, but we pick up on a few issues for improvement of bio-aerosol MS in the next few years.

For bio-aerosol analysis, pyMS is currently best developed among the aerosol MS technologies. Given the fact that, so far, pyGC-ion mobility spectrometry (IMS) was developed for field bio-aerosol detection (Snyder et al. 2004), the technology and chemistry of FAME pyMS can be further developed for bio-aerosol detection (Krebs et al. 2006; Prasad et al. 2006). Changing from MS to IMS technology does resolve design issues with regard to size and transportability of the overall system. Although pyMS has the advantage that it does not include relatively vulnerable lasers, the use of laser ionization may greatly aid the selectivity in ionization of specific pyrolysis products. For example, photoionization was demonstrated for the analysis of polymers by pyMS (see, e.g., Zoller et al. 1999), for the field analysis of chemical weapons agents (Syage, Hanning-Lee, and Hanold 2000), and for the direct analysis of biological material (Evans, Hanold, and Syage 2000; Nies, Evans, and Syage 2003). Alternatively, surface ionization might give useful results when applied to bio-aerosols. Surface ionization is already employed in aerosol MS (e.g., Jimenez et al. 2003), but the published applications were limited to organic and inorganic aerosols. Simplification of the instrument by going from MS to IMS and application of other ionization modes, such as photoionization or surface ionization, may provide new signatures for the analysis of bio-aerosol components.

For all bio-aerosol MS technologies, downsizing is an important issue on the way to increased transportability. Downsizing applies to all of the component technologies integrated in a bio-aerosol mass spectrometer. Downsizing of the mass spectrometer has the additional
benefit that smaller vacuum volumes require less pumping capacity and, hence, smaller pumps. The application of Ion Trap MS instead of TOF MS will sometimes be an option in downsizing, and we see Ion Trap MS (e.g., Harris, Reilly, and Whitten 2006) as a viable line of development in bio-aerosol MS. However, both types of mass spectrometer have proved amenable to downsizing, as has been reviewed by Badman and Cooks (2000). Downsizing often involves application of different or refined physical principles, with a concomitant redesign of the mass analyzer. The application of higher order ion focusing and a redesign of the instrument has for example produced small “end cap” reflectron TOF instruments with capabilities comparable to those of common laboratory TOF instruments (e.g., Cornish and Cotter 1997; Fancher, Woods, and Cotter 2000). A completely different way was pursued with the redesign of the reflectron from the common multistage to a single stage element (Uphoff, Muskat, and Grotemeyer 2004). Also, developments in the manufacturing process have led to smaller TOF instruments, even down to suitcase size (Cornish, Ecelberger, and Brinckerhoff 2000; Ecelberger et al. 2004). Downsizing of Ion Traps to a hand-portable size was recently accomplished (Song et al. 2006; Gao et al. 2006), although it should be noted that the mass range of such small Ion Traps is not yet compatible with high mass range applications. For either TOF or Ion Trap instruments, downsizing of the mass spectrometer will require tailoring to the specific application. Downsizing of laser systems will also drastically reduce the system size of LDI or MALDI bio-aerosol MS systems, with the additional benefit that smaller lasers require less cooling. The development of lasers is mainly driven by other applications, but more specific requirements for (aerosol) mass spectrometry have recently been pointed out in great detail (Holle et al. 2006; Murphy 2006). In particular, the development of solid-state lasers with UV capability will help in downsizing bio-aerosol MS systems. Downsizing of laser equipment and mass analyzers will determine the overall system downsizing, because aerosol sampling and capture equipment will not be amenable to much downsizing given the required performance.

For all bio-aerosol technologies, data handling is of the essence. Spectrum comparison with library spectra is a routine application, but actual identification of bacteria from spectra requires more research. For any of the bio-aerosol analysis methods applied to bacteria, robustness of the signature towards environmental factors and phenotype requires more research. The signatures obtained from pyMS and LDI approaches will not have sufficient distinctive capability for adequate identification, because the ionized material is essentially a far derivative of biological material. Nevertheless, in the analysis of bacterial bio-aerosols, pyMS and LDI signatures may be useful as indicators rather than identifiers. As for the MALDI MS approach to bio-aerosol analysis, the obtained signatures relate directly to bacterial biomolecules present. However, more research is required to establish the link between aerosol MALDI MS signature and bacterial biomolecules. On the basis of the current state of the art in laboratory MALDI MS of bacteria, direct aerosol MALDI MS has the potential to become a rapid identification method.

In conclusion, bio-aerosol MS still poses a challenge to science and technology, with interesting promises of environmental, medical and safety applications to be fulfilled in the next few years.

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