A commentary on the role of molecular technology and automation in clinical diagnostics

Ciara O’Connor¹,², Marie Fitzgibbon¹, James Powell¹, Denis Barron¹, Jim O’Mahony¹, Lorraine Power¹, Nuala H O’Connell¹,², and Colum Dunne²,∗

¹Department of Clinical Microbiology; University Hospital Limerick; Limerick, Ireland; ²Centre for Interventions in Infection, Inflammation, and Immunity (4i) and Graduate Entry Medical School; University of Limerick; Limerick, Ireland; ³Cork Institute of Technology; Cork, Ireland

Keywords: clinical microbiology, MALDI, PCR, patient care, impact

Historically, the identification of bacterial or yeast isolates has been based on phenotypic characteristics such as growth on defined media, colony morphology, Gram stain, and various biochemical reactions, with significant delay in diagnosis. Clinical microbiology as a medical specialty has embraced advances in molecular technology for rapid species identification with broad-range 16S rDNA polymerase chain reaction (PCR) and matrix-assisted laser desorption and/or ionization time of flight (MALDI-TOF) mass spectrometry demonstrated as accurate, rapid, and cost-effective methods for the identification of most, but not all, bacteria and yeasts. Protracted conventional incubation times previously necessary to identify certain species have been mitigated, affording patients quicker diagnosis with associated reduction in exposure to empiric broad-spectrum antimicrobial therapy and shortened hospital stay. This short commentary details such molecular advances and their implications in the clinical microbiology setting.

Introduction

Europe and North America have experienced significant declines in mortality secondary to infectious diseases.⁵ Indeed, the incidence of severe sepsis, associated with multiorgan failure, in the European Union is currently estimated as 90.4 cases per 100,000 population.² Global developments in medicine, including advances in education and training for medical students, continuous professional development for qualified physicians, modern cleaner hospitals with single en-suite accommodation, improved infection control practices, newer antimicrobial therapies, advanced molecular technology used in the laboratory, and sepsis management protocols have all contributed to the decline in sepsis-related deaths.⁵ Previous studies have shown a direct link between outcomes from infectious illnesses and time to pathogen identification.⁴ Consequently, laboratory testing volumes are increasing by 10–15% per year internationally, driven partly by infection control demands, with enhanced screening for methicillin-resistant Staphylococcus aureus and multi-drug resistant organisms such as vancomycin-resistant enterococci, extended spectrum β-lactamases, and carbapenemase-resistant enterobacteriaceae.⁵

Clinical microbiology has evolved in response to clinical needs and laboratories have adapted to meet demand for testing. The laboratory service expected by physicians and patients has changed dramatically. Most Western tertiary referral hospitals operate a laboratory service capable of providing an on-call 24 hour service enabling clinical teams to obtain results as soon they become available. Patients, many empowered and self-educated through the vast array of information readily available on the Internet, expect rapid reporting of results with minimal delay from time of presentation to diagnosis and discharge. Patients are also increasingly aware of the dangers of acquiring nosocomial illnesses, and there are also many added economic and financial pressures being brought to bear on clinical services. As a consequence, approximately 80% of patient management decisions are influenced by laboratory testing⁶ and reducing time to identification within the laboratory is increasingly a priority.

The Complementary Roles of the Clinical Microbiologist and the Microbiology Laboratory

The primary roles of the clinical microbiology team are to guide and support physicians in community or hospital settings, to select appropriate diagnostic investigations and antimicrobials, as warranted, and to achieve the best possible outcome for patients. For any patient, an array of samples may be sent for laboratory analysis with the objective of identifying causative pathogens. Specimens may be analyzed following sampling from diverse physiological sources including, but not limited to, cerebro-spinal fluid, blood, “sterile” body fluids, tissue and pus, urine, intravascular cathether tips, prosthetic devices, and the respiratory tract. Equipped with expertise and experience, clinical microbiologists are responsible for construction of differential diagnoses and provision of advice on required testing. However, skilled medical scientists are critical to the processes of appropriate growth media selection, inoculation
and incubation of specimens, and analysis and interpretation of complex analytical data.

**Patient-Centered Care**

Patients presenting for medical attention with signs of infection, particularly evidence of fever, with or without hemodynamic instability, should have at least one set of blood cultures taken using an appropriate aseptic technique. Blood samples forwarded by physicians to the laboratory, that flag as positive when incubated, are typically the first engagement of clinical microbiologists with patients who may require either commencement or escalation of antimicrobial therapy. However, blood cultures are problematic in diagnosis of sepsis due to the potential presence of unculturable organisms, underfilling of blood collection vessels, protracted incubation times, and false positives caused by leucocytosis or contamination with commensal flora (e.g., coagulase negative staphylococci). In addition, microorganism detection and susceptibility testing may require up to 48 hours, delaying definitive diagnoses for patients.

**Why the Long Delay?**

Processing of microbiology samples can be a lengthy process (Fig. 1). Once a blood sample is confirmed as positive in the laboratory, preliminary organism identification—usually performed by medical scientists—is determined by Gram stain, traditional culture techniques, and colony morphology. Microscopic examination of stained smears of tissue or biological fluid is relatively quick and requires minimal resource inputs. However, such assessment may have low sensitivity and specificity as, historically it has been grounded in subjective criteria such as odor, color, and experience of phenotypic patterns derived using a variety of confirmatory biochemical tests performed at the laboratory bench. Subsequent, authorized clinical diagnosis is reliant on the organism being culturable on solid media following 24 hours (aerobic) or 48 hours (anaerobic) incubation. Frequently, incubation can prolong the process by 24–48 hours.

The duration of processes involved in reaching definitive laboratory diagnoses via these conventional culture methods results in widespread utilization of empiric broad-spectrum antimicrobial therapy in stabilizing of deteriorating patients. Clearly, the intention is to administer a broad spectrum of bacteriostatic and/or bacteriocidal agents sufficient to treat the unidentified pathogen, but, in reality, this approach is often associated with development of antimicrobial resistance, *Clostridium difficile* overgrowth, and iatrogenic complications for patients.7,8 Typically, empiric antimicrobial therapy is rationalized to a narrower spectrum agent once microbiology laboratory results become available.

**Can Laboratory Results Be Expedited?**

Matrix-assisted laser desorption ionisation–time of flight mass spectrometry (MALDI-TOF MS)9-11 has facilitated substantial movement toward same-day clinical decision-making underpinned by molecular microbiology (Fig. 2).12 In use, MALDI-TOF MS determines characteristic protein patterns derived from microbe composition and allows interrogation of well-developed databases for identification. This can occur once an incubated sample grows to colony level (approximately 10⁵ cfu) and can be transferred to a slide. Laser analysis then takes less than 10 min, significantly reducing organism identification times compared with conventional microbiology.

Increasingly utilized clinically for identification of bacteria,8,13,14 and yeasts,15 MALDI-TOF MS has potential for accurate identification of viruses.16 Its utilization in clinical laboratories has resulted in reports of previously undetected organisms, such as new species of anaerobes17,18 (e.g., Prevotella spp. and Anaerococcus spp.). However, while MALDI-TOF MS has a reported reliability of >95% (of routine isolates grown on solid media in the laboratory),19 its limitations include inability to differentiate genetically similar organisms, such as *Streptococcus pneumoniae* from other members of the *Streptococcus mitis*.
group, and Shigella species from Escherichia coli. Despite this, MALDI-TOF MS analysis can reduce manual workload for medical scientists and improve throughput of laboratory samples. Exemplifying this, previously technically-challenging identification of microorganisms from sputum of cystic fibrosis patients, utilizing multiple selective media and prolonged incubation, may now be completed within 48 hours of sampling. More specifically, a 2012 Canadian study reported a mean reduction in turnaround time of 34.3 hours when definitive identification of isolates directly from blood samples was possible, at 26.5 hours when incubation was necessary before MALDI-TOF MS could be utilized. These advances are especially relevant clinically as identification of Gram-negative and Gram-positive bacteria as well as yeasts can now be obtained directly from 7–8 ml of blood when using systems such as the MALDI Sepsityper® Kit (BrukerDaltonics 2010), thereby eliminating incubation times associated with growth on solid media.

In an attempt to replicate the reductions observed in Canada (described above), 150 blood samples, flagged as positive for presence of microorganisms using the BacT/ALERT® system (bioMérieux®), were analyzed at University Hospital Limerick (UHL) over a 6 week period using the MALDI Sepsityper® Kit and conventional microbiological techniques, and results compared with respect to turnaround time (TAT) of positive blood cultures (Table 1). We observed that for polymicrobial blood cultures, MALDI-TOF resulted in TAT being reduced by a mean of 17.5 hours, mean reduction of 14.47 hours for Gram-positive cultures and mean reduction of 23.19 hours for Gram-negative cultures.

The impact of our study was that, of the patients from whom these blood samples were taken, 15 patients (30.0%) experienced a change of therapy in a timelier manner, 12 patients (24%) were changed to a more appropriate antibiotic, while one patient (2%) had antibiotic therapy discontinued. These changes correspond with findings elsewhere that rapid detection of microbes facilitates the introduction of appropriate narrow spectrum targeted therapy and reduction of patient length of stay or, indeed, avoidance altogether of admission via outpatient antibiotic therapy (OPAT).

Further, MALDI-TOF is both cost and environmentally effective when compared with conventional microbiology. Gaillot et al. demonstrated 89.3% cost savings in the first year of MALDI-TOF use, whereby cost per isolate decreased from $5.80 to $0.50 and waste reduced from >1400 kg to <50 kg, in addition to lesser need for DNA and/or RNA sequencing and, overall, significantly enhanced time to bacterial identification. At the time of writing, microbiological testing at UHL was predominantly based on the bioMérieux® ARIS system costing approximately €3.50 per identification, and with ca. €17000 annual expenditure on supplementary identification methods (bioMérieux®API® and RemelRapID®).

**PCR and Patients**

Polymerase chain reaction (PCR) is used extensively for identification of multiple pathogens, commonly including methicillin-resistant Staphylococcus aureus, Clostridium difficile, Streptococcus agalactiae (group B Streptococcus), and bacterial causes of meningitis. At UHL, polymerase chain reaction (PCR) is used extensively in testing for Chlamydia trachomatis and Neisseria gonorrhoeae. Between January and December 2013, 13338 clinical samples, on average 1100 per month, were analyzed using PCR (Table 2). Such testing introduced a “same-day service,” with results available for clinicians within hours of a patient’s attendance at an on-site sexual health clinic, facilitating early commencement of directed-therapy and reduced patient anxiety. We also use PCR in evaluating success or otherwise of
antiviral therapies and for long-term follow-up of patients with HIV and Hepatitis B and C attending our local services.

Decision-making regarding respiratory viruses, for example, respiratory syncitial virus (RSV) in pediatric patients has been reduced at UHL from up to a week to just hours, with significant benefits for hospital bed management and prompt initiation of antiviral therapy as necessary, and as reported elsewhere. Detection of Herpes viruses and a variety of enteroviruses via PCR is performed on all cerebrospinal fluid samples received in the laboratory, thus supporting appropriate timely management of common viral central nervous system infections.

In our hands, PCR for analysis of stool samples has reduced turnaround time for samples by up to 24 h. C. difficile toxin testing via PCR was introduced at UHL in September 2013, with over 400 stool samples now analyzed each month. At a practical level, given the transmissibility of C. difficile, the need for urgent isolation, commencement of treatment, and the potential for serious complications such as toxic megacolon, the introduction of PCR has considerably improved our patient service.

PCR technology continues to evolve and ready analysis of samples obtained directly from clinical specimens such as pleural fluid, central spinal fluid, blood, joint aspirates, heart valves, and abscess aspirates is increasingly commonplace. PCR technology continues to evolve and ready analysis of samples obtained directly from clinical specimens such as pleural fluid, central spinal fluid, blood, joint aspirates, heart valves, and abscess aspirates is increasingly commonplace. PCR technology continues to evolve and ready analysis of samples obtained directly from clinical specimens such as pleural fluid, central spinal fluid, blood, joint aspirates, heart valves, and abscess aspirates is increasingly commonplace. Regularly, broad-range 16S rDNA PCR is utilized in the context of conventional microbiology has not confirmed infection.

### Table 1. Turnaround time (TAT) for microbial identification

|                             | MALDI-TOF | Conventional microbiology | Reduction of mean TAT |
|-----------------------------|-----------|---------------------------|-----------------------|
|                             | Mean      | Min- Max                  | Mean                  | Min- Max |
| Gram-positive and Gram-negative | 11.12***  | 2.0–27                    | 28.62                 | 18.0–60.0 | 17.5     |
| Gram-positive only           | 13.41     | 2.0–27.0                  | 27.88                 | 20.0–55.0 | 14.47    |
| Gram-negative only           | 5.36      | 2.0–20.0                  | 28.55                 | 18.0–60.0 | 23.19    |

*TAT = turnaround time. **Over a 6 wk period, 150 blood samples at University Hospital Limerick, Ireland. ***All values are in hours.

### Table 2. PCR analysis at University Hospital Limerick

|                             | Annual | Monthly |
|-----------------------------|--------|---------|
|                             | Mean   | Min- Max |
| Total samples tested        | 15732  | 1311    | 2.0–27 |
| Samples tested for potential STI only | 13338  | 1112    | 769–1148 |
| For *Clostridium difficile* (toxin) | 1714d  | 429     | 415–447 |
| For *Salmonella*, *Shigella*, *Campylobacter*, VTEC | 2680e  | 670     | 544–941 |

*January–December 2013; discrete clinical samples tested. STI: sexually transmitted infection, specifically *Chlamydia trachomatis* and *Neisseria gonorrhoeae*. Introduced at UHL in September 2013. Projected annual total based on 4 mo analysis: 5142. Projected annual total based on 4 mo analysis: 8040.

Practical Issues Associated with Molecular Microbiology

No diagnostic technology is universally applicable. For example, important pathogens such as *Streptococcus pneumoniae*, β-hemolytic streptococci, and enteric pathogens such as *Shigella spp.*, and *Escherichia coli* remain difficult to identify using MALDI-TOF MS. For this reason, microbiologists remain reliant on traditional methods of identification and for antimicrobial susceptibility testing. On a financial note, there are also capital funding considerations in adopting molecular technologies. Many clinical laboratories will require refurbishment, or new builds, before being suitable to accommodate molecular equipment such as MALDI-TOF MS or to facilitate laboratory automation using suites such as Becton Dickinson’s Kiestra™. Indeed, movement toward sophistication of laboratory methods should ideally to be integrated with electronic patient records (i.e., [@national unique patient identification numbers]) to avoid duplication of unnecessary tests caused by patients having duplicate or triplicate patient laboratory registration numbers. At least in Ireland, such a system does not yet exist. An equally important consideration is the requirement for upskilling of staff and maintenance of expensive equipment in the context of stringent laboratory accreditation requirements, such as compliance with ISO standard 17025.

### Training in the Molecular Era

There are challenges for clinical microbiologists and physicians in keeping pace with rapidly changing discoveries in molecular diagnostic techniques, for instance, the potential use of MALDI-TOF or high-resolution melting PCR analysis in assessment of antimicrobial susceptibility and isolate profiling. Training programs for specialist physicians in clinical microbiology will need to take greater account of this and allow time for medical professionals to become and remain familiar with emerging developments, both their advantages and, from a patient safety perspective, their limitations. A further risk to be mitigated is the potential de-skilling of medical scientists with the advent of, and over-reliance on, modern technologies.
Conclusions

Clinical microbiology is pivotal to patient outcomes in providing highly accurate diagnostic and supportive services to clinicians across all specialties in acute settings and primary care. The introduction of molecular technology and automation fundamentally changes the way in which laboratory diagnoses are reached, providing superior laboratory services to physicians and patients. Molecular diagnostics afford improved sensitivity and specificity, and the possibility of rapid diagnoses, with turnaround times of hours rather than days, with subsequent reduced length of hospital stay. The increase in antimicrobial resistance globally reinforces the need for more rapid diagnostics, facilitating judicious use of effective antimicrobials and mitigated risk of hospital-acquired infections and, in summary, improving patient-centered care.

Disclosure of Potential Conflicts of Interest

No potential conflict of interest was disclosed.

Acknowledgments

We are grateful to the staff of the microbiology laboratory, and to Mike O’Nolan (IT Department), at University Hospital Limerick for their contributions to this commentary.

References

1. Armstrong GL, Conn LA, Pinner RW. Trends in infectious disease mortality in the United States during the 20th century. JAMA 1999; 281:61-6; PMID:9892452; http://dx.doi.org/10.1001/jama.281.1.61
2. Daniels R. Surviving the first few hours in sepsis: getting the basics right (an intensivist’s perspective). J Antimicrob Chemother 2011; 66(Suppl 2):iii1-23; PMID:21398303; http://dx.doi.org/10.1093/jac/dkt515
3. Phua J, Ho BC, Tee A, Chan KP, Johan A, Loo S, So CR, Chia N, Tan AY, Tham HM, et al. The impact of clinical protocols in the management of severe sepsis: a prospective cohort study. Anaesth Intensive Care 2012; 40:663-74; PMID:22815395
4. Yang S, Rothman RE. PCR-based diagnostics for infectious diseases: uses, limitations, and future applications in acute-care settings. Lancet Infect Dis 2004; 4:337-48; PMID:15172342; http://dx.doi.org/10.1016/S1473-3099(04)01044-8
5. Bourbeau PP, Leboeber NA. Automation in clinical microbiology. J Clin Microbiol 2013; 51:1658-65; PMID:23555547; http://dx.doi.org/10.1128/JCM.00301-13
6. Dokouhaki P, Blondeau JM. Advances in laboratory diagnostic technologies in clinical microbiology and what this means for clinical practice. Clin Pract 2012; 9:347-52; http://dx.doi.org/10.2217/cpr.12.32
7. Dusywna W. Strategies of empiric antibiotic therapy in severe sepsis. Anaesthesiol Intens Ther 2012; 44:96-105; PMID:22992970
8. Carleton A, Casserly B, Power L, Linnane B, O’Laherry G, Powell J, Harrnett P, Collins J, Murphy P, Kenna D, et al. Clustered multidrug-resistant Bordetella petrii in adult cystic fibrosis patients in Ireland: case report and review of antimicrobial therapies. JMM Case Reports 2014; 1:1-6
9. Carbonnelle E, Mesquita C, Bille E, Day N, Dauphin B, Beretti JL, Ferroni A, Gutmann L, Nassif X. MALDI-TOF mass spectrometry tools for bacterial identification in clinical microbiology laboratory. Clin Biochem 2011; 44:104-9; PMID:20862154; http://dx.doi.org/10.1016/jClinBiochem.2010.06.017
10. Seng P, Rolain JM, Fournier PE, La Scola B, Drancourt M, Raoult D. MALDI-TOF mass spectrometry applications in clinical microbiology. Future Microbiol 2010; 5:1733-54; PMID:21136362; http://dx.doi.org/10.2217/fmb.10.127
11. Neville SA, Lecordier A, Zechos H, Chater MJ, Gosbell IB, Maley MW, van Hal SJ. Utility of matrix-assisted laser desorption ionisation-time of flight mass spectrometry following introduction for routine laboratory bacterial identification. J Clin Microbiol 2011; 49:2980-4; PMID:21632894; http://dx.doi.org/10.1128/JCM.00438-11
12. Bullman S, Lucey B, Seator RD. Molecular diagnostics: the changing culture of medical microbiology. Bioeng Bugs 2012; 3:1-7; PMID:22279143; http://dx.doi.org/10.4161/bbi.3.1.19011
13. Bizzini A, Durussel C, Bille J, Greub G, Prod’hom G. Performance of matrix-assisted laser desorption ionization-time of flight mass spectrometry for identification of bacterial strains routinely isolated in a clinical microbiology laboratory. J Clin Microbiol 2010; 48:1549-54; PMID:20202166; http://dx.doi.org/10.1128/JCM.00794-09
14. Loonen AJ, Janz AR, Berglund JN, Valkenburg M, Wolk DM, van den Brule AJ. Comparative study using phenotypic, genotypic, and proteomics methods for identification of coagulate-negative staphylococci. J Clin Microbiol 2012; 50:1437-9; PMID:22238435; http://dx.doi.org/10.1128/JCM.00674-11
15. Santos C, Paterson RR, Venâncio A, Lima N, Filamentoung fungal characterizations by matrix-assisted laser desorption/ionization-time of flight mass spectrometry. J Appl Microbiol 2010; 108:375-85; PMID:19659699; http://dx.doi.org/10.1111/j.1365-2672.2009.04448.x
16. Cobo F. Application of maldi-tof mass spectrometry in clinical virology: a review. Open Virol J 2013; 7:78-90; PMID:24222805; http://dx.doi.org/10.2174/18743579201307200927003
17. La Scola B, Fournier PE, Raoult D. Burden of nonpneumococcal streptococci of the Streptococcus mitis group by matrix-assisted laser desorption/ionization-time of flight mass spectrometry. J Clin Microbiol 2011; 49:1437-9; PMID:21762636; http://dx.doi.org/10.1128/JCM.01794-09
18. Barrea M, Pagnier I, La Scola B. Improving the identification of anaerobes in the MALDI-TOF and 16S rRNA gene sequencing era. Anaerobe 2011; 17:106-12; PMID:21672636; http://dx.doi.org/10.1016/j.anaerobe.2010.05.010
19. Barreau M, Pagnier I, La Scola B. Improving the identification of anaerobes in the clinical microbiology laboratory through MALDI-TOF mass spectrometry. Anaerobe 2013; 22:123-5; PMID:23639482; http://dx.doi.org/10.1016/j.anaerobe.2013.04.011
20. Carbonnelle E, Gros P, Jacquier H, Day N, Tenza S, Dewailly A, Vissouarn O, Rottman M, Herrmann KA, Pang PP, Guenterer J, Webb AA, Miller C, Alfa MJ. Identification of blood culture isolates directly from positive blood cultures by use of matrix-assisted laser desorption ionization-time of flight mass spectrometry and a commercial extraction system: analysis of performance, cost, and turnaround time. J Clin Microbiol 2012; 50:3324-8; PMID:22875888; http://dx.doi.org/10.1128/JCM.01479-12
21. Schieffer KM, Tan KE, Stamper PD, Somogyi A, Andrea SB, Wakefield T, Romagnoli M, Chapin KC, Wolk DM, Carroll KC. Multicenter evaluation of the Sequipart™ extraction kit and MALDI-TOF MS for direct identification of positive blood culture isolates using the BD BACTECTM FX and VersaTREK® diagnostic blood culture systems. J Appl Microbiol 2014; 116:954-41; PMID:24410849; http://dx.doi.org/10.1111/jam.12434
22. Tenover FC. Potential impact of rapid diagnostic tests on improving antimicrobial use. Ann N Y Acad Sci 2010; 1213:70-80; PMID:20514433; http://dx.doi.org/10.1111/j.1749-6632.2010.05827.x
23. Gauthier O, Blondiaux N, Loize C, Waller F, Lemaître N, Herwegh S, Courcol RJ. Cost-effectiveness of switch to matrix-assisted laser desorption ionization-time of flight mass spectrometry for routine bacterial identification. J Clin Microbiol 2011; 49:4412; PMID:21998417; http://dx.doi.org/10.1128/JCM.05429-11
24. Steed LL, Ball RT Jr. Respiratory virus detection: beyond influenza and RSV into emerging infectious diseases. J S C Med Assoc 2013; 109:85-7; PMID:23650422; http://dx.doi.org/10.1128/JCM.00438-11

©2014 Landes Bioscience. Do not distribute.
29. López Roa P, Alonso R, de Egea V, Usubillaga R, Muñoz P, Bouza E. PCR for detection of herpes simplex virus in cerebrospinal fluid: alternative acceptance criteria for diagnostic workup. J Clin Microbiol 2013; 51:2880-3; PMID:23804382; http://dx.doi.org/10.1128/JCM.00950-13

30. Jensen KH, Dargis R, Christensen JJ, Kemp M. Ribosomal PCR and DNA sequencing for detection and identification of bacteria: experience from 6 years of routine analyses of patient samples. APMIS 2014; 122:248-55; PMID:23879657; http://dx.doi.org/10.1111/apm.12139

31. Duffert S, Missaghi B, Daley P. Culture-negative endocarditis diagnosed using 16S DNA polymerase chain reaction. Can J Infect Dis Med Microbiol 2012; 23:216-8; PMID:24294278

32. Kostrzewa M, Spahier K, Maier T, Schubert S. MALDI-TOF MS: an upcoming tool for rapid detection of antibiotic resistance in microorganisms. Proteomics Clin Appl 2013; 7:767-78; PMID:24123965; http://dx.doi.org/10.1002/prca.201300042

33. Gabriel EM, Douarre PE, Fitzgibbon S, Clair J, Lucey B, Coffey A, O’Mahony JM. High-resolution melting analysis for rapid detection of linezolid resistance (mediated by G2576T mutation) in Staphylococcus epidermidis. J Microbiol Methods 2012; 90:134-6; PMID:22561095; http://dx.doi.org/10.1016/j.mimet.2012.04.002