Mycolactone diffuses into the peripheral blood of Buruli ulcer patients—implications for diagnosis and disease monitoring.

Fred S. Sarfo, Fabien Le Chevalier, N’Guetta Aka, Richard O. Phillips, Yaw Amoako, Ivo G. Boneca, Pascal Lenormand, Mireille Dosso, Mark Wansbrough-Jones, Romain Veyron-Churlet, et al.

To cite this version:
Fred S. Sarfo, Fabien Le Chevalier, N’Guetta Aka, Richard O. Phillips, Yaw Amoako, et al.. Mycolactone diffuses into the peripheral blood of Buruli ulcer patients—implications for diagnosis and disease monitoring.. PLoS Neglected Tropical Diseases, 2011, 5 (7), pp.e1237. 10.1371/journal.pntd.0001237 . inserm-00691438

HAL Id: inserm-00691438
https://www.hal.inserm.fr/inserm-00691438
Submitted on 26 Apr 2012

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Mycolactone Diffuses into the Peripheral Blood of Buruli Ulcer Patients - Implications for Diagnosis and Disease Monitoring

Fred S. Sarfo1,*, Fabien Le Chevalier2,3, N’Guetta Aka3, Richard O. Phillips1,4, Yaw Amoako1, Ivo G. Boneca5,6, Pascal Lenormand7, Mireille Dosso3, Mark Wansbrough-Jones8, Romain Veyron-Churlet2, Laure Guenin-Macé2, Caroline Demangel2

1 Komfo Anokye Teaching Hospital, Kumasi, Ghana, 2 Institut Pasteur, Pathogénomique Mycobactérienne Intégrée, Paris, France, 3 Institut Pasteur, Mycobactéries Tuberculeuses et Atypiques, Abidjan, Côte d’Ivoire, 4 Kwame Nkrumah University of Science and Technology, Kumasi, Ghana, 5 Institut Pasteur, Biologie et Génétique de la Paroi Bactérienne, Paris, France, 6 INSERM, Groupe AVENIR, Paris, France, 7 Institut Pasteur, Plateforme Protéomique, Paris, France, 8 St George’s University of London, London, United Kingdom

Abstract

Background: Mycobacterium ulcerans, the causative agent of Buruli ulcer (BU), is unique among human pathogens in its capacity to produce a polyketide-derived macrolide called mycolactone, making this molecule an attractive candidate target for diagnosis and disease monitoring. Whether mycolactone diffuses from ulcerated lesions in clinically accessible samples and is modulated by antibiotic therapy remained to be established.

Methodology/Principal Finding: Peripheral blood and ulcer exudates were sampled from patients at various stages of antibiotic therapy in Ghana and Ivory Coast. Total lipids were extracted from serum, white cell pellets and ulcer exudates with organic solvents. The presence of mycolactone in these extracts was then analyzed by a recently published, field-friendly method using thin layer chromatography and fluorescence detection. This approach did not allow us to detect mycolactone accurately, because of a high background due to co-extracted human lipids. We thus used a previously established approach based on high performance liquid chromatography coupled to mass spectrometry. By this means, we could identify structurally intact mycolactone in ulcer exudates and serum of patients, and evaluate the impact of antibiotic treatment on the concentration of mycolactone.

Conclusions/Significance: Our study provides the proof of concept that assays based on mycolactone detection in serum and ulcer exudates can form the basis of BU diagnostic tests. However, the identification of mycolactone required a technology that is not compatible with field conditions and point-of-care assays for mycolactone detection remain to be worked out. Notably, we found mycolactone in ulcer exudates harvested at the end of antibiotic therapy, suggesting that the toxin is eliminated by BU patients at a slow rate. Our results also indicated that mycolactone titres in the serum may reflect a positive response to antibiotics, a possibility that it will be interesting to examine further through longitudinal studies.

Introduction

Buruli ulcer (BU), caused by Mycobacterium ulcerans, is the third most common mycobacterial disease after tuberculosis and leprosy and represents an emerging threat [1,2]. Since the late 1980s, the disease has been developing throughout West and Central Africa, prompting the WHO in 1998 to initiate an awareness and control campaign (http://www.who.int/gtb-buruli). Although efficient [3,4,5,6,7], current treatment protocols recommend the daily administration of oral rifampicin and intramuscular streptomycin for 8 weeks, with additional surgical intervention when necessary. To control the emergence of BU and to improve the management of the disease, it is vital to develop new tools for early diagnosis and treatment monitoring. A distinctive feature of M. ulcerans among human pathogens is the production of mycolactone [8], a macrocyclic polyketide playing a critical role in bacterial virulence (reviewed in [9,10]). We have recently demonstrated the presence of intact mycolactone in punch biopsies from all forms of BU disease, before and during antibiotic therapy [11]. Moreover, there is evidence from mouse studies that mycolactone may diffuse into the peripheral blood.
12]. Here we used chemical approaches to determine if mycolactone is present in blood samples and ulcer exudates obtained non-invasively at various stages of antibiotic therapy.

Methods

Patient cohorts

Patients were recruited if they met the WHO clinical case definition of BU disease; were not pregnant; had no history of tuberculosis, leprosy, or liver, kidney, or hearing impairment. All subjects provided written informed consent (thumb print of parent or guardian in the case of children, depending on literacy). A cross-section of patients with BU disease were recruited which included a spectrum of patients yet to initiate antibiotic therapy, some at various stages of antibiotic treatment and few who had completed treatment. Healthy controls from the same endemic area were included. In Ghana, patients were recruited by local health workers from villages near Tepa Government Hospital in the Ahfo Ano North District of Ghana, where there is a high prevalence of BU. The study protocol was approved by the ethics review committees at the School of Medical Sciences, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana. In Ivory Coast, patients were either recruited from the Djekanou General Hospital or detected by a mobile medical team actively screening the district of Abidjan. The study protocol was approved by the national ethic review committee.

Diagnosis and treatment

To confirm the clinical diagnosis, punch biopsy specimens of 4-mm diameter (Ghana) or ulcer exudates (Ivory Coast) were tested by PCR for the IS2404 repeat sequence, which is characteristic of M. ulcerans. Positive patients were treated with 10 mg/kg oral rifampicin and 15 mg/kg intramuscular streptomycin daily, administered at village health posts under direct observation, according to the WHO recommendations. Only IS2404 positive samples were considered for analysis of mycolactone presence.

Mycolactone standard

Mycolactone was extracted from M. ulcerans 1615 (ATCC 35840), as previously described [12]. In brief, bacteria were cultivated in Middlebrook 7H9 broth (Difco) enriched with 10% oleic acid-albumin-dextrose-catalase (OADC, Becton Dickinson) for 4 weeks in spinner flasks at 30°C. Total lipids were extracted from bacterial cell pellets with 2/1 CHCl3/MeOH (v/v) for 20 h at 4°C. After separation from the aqueous phase following the addition of 20% H2O (v/v), the organic phase was dried. The resulting material was resuspended in ice-cold acetone and

Figure 1. Comparison of the performances of the TLC-Fluo and HPLC approaches for mycolactone quantitative detection. A representative picture of the fluorescent signals and elution peaks obtained for mycolactone (10–500 ng) detection by TLC-Fluo (A) and HPLC (B) are shown, with the corresponding standard curves (C). Arbitrary units correspond to fluorescence intensity of the mycolactone band (TLC-Fluo) and area of the mycolactone elution peak (HPLC). Similar results were obtained in at least three independent experiments.

doi:10.1371/journal.pntd.0001237.g001
incubated for 20 h at −20°C. The acetone-soluble fraction was then dried, resuspended in ethanol, and loaded onto a silica gel TLC plate and eluted with 90/9/1 CHCl₃/MeOH/H₂O as the mobile phase. The yellow band corresponding to mycolactone (retention factor of 0.2) was then scraped and mycolactone eluted from silica particles using 2/1 CHCl₃/MeOH (v/v). Following solvent evaporation, purified mycolactone was resuspended in ethanol. The concentration of the resulting solution was determined by UV absorption, as described [13].

Mycolactone detection by coupling to 2-naphtalene boronic acid (TLC-Fluo)

As recently described [13], mycolactone or lipid extracts were applied to a silica gel TLC plate and eluted with 90/9/1 CHCl₃/MeOH/H₂O as the mobile phase. The eluted TLC plate was briefly warmed on a hot plate to evaporate the organic solvents, and quickly immersed into a 0.1 M acetone solution of 2-naphtalene boronic acid (Sigma), then heated to 100°C for 5–10 seconds. The TLC plate was then irradiated with a UV reader equipped with a 312 nm lamp.

Mycolactone detection by HPLC/MS/MS

Mycolactone or lipid extracts were also analyzed by High Performance Liquid Chromatography coupled to Mass Spectrometry (HPLC/MS/MS). We used a Shimadzu HPLC fitted with a BDS Hypersil C8 column (5 µm, 4.6×250 mm), with UV detection at 360 nm. Mycolactone was eluted by a 60 min gradient from 50 to 95% acetonitrile in water after 22 min. Fractions of interest were collected in glass tubes and analysed on a QSTAR XL, AB-MDS-SCIEX mass spectrometer with an electrospray ion source and the following parameters: ion spray voltage (IS), 5200 v; curtain gas (CUR), 25; gas 1 (GS1), 5;

Figure 2. Structurally intact mycolactone is detected in ulcer exudates of BU patients. (A) Example of the fluorescence signals given by ulcer exudates (lanes 2–5), compared to pure mycolactone (lane 1) following analysis by TLC-Fluo. The band corresponding to mycolactone is designated by an asterisk. (B) HPLC elution profiles are shown for reference mycolactone (100 ng) and for one representative ulcer exudate among the 20 positive ones. The corresponding MS/MS spectra are presented, with mycolactone parent ion (P.I) and products designated by asterisks. Similar MS/MS spectra were obtained from HPLC elution peaks collected from three positive patients.

doi:10.1371/journal.pntd.0001237.g002
declustering potential (DP), 50 v; focusing potential (FP), 225 v; declustering potential 2 (DP2), 15 v. Mycolactone was identified by the presence of [M+Na]+ m/z 765.5. MS-MS parameters were: ion spray voltage (IS), 6000 v; curtain gas (CUR), 25; gas 1 (GS1), 20; declustering potential (DP), 60 v; focusing potential (FP), 230 v; declustering potential 2 (DP2), 10 v; collision energy (CE), 60; collision gas (CAD), 5. Data were collected and processed through the Analyst QS 1.1 software from AB-MDS-Sciex.

Exudates
Wound swabs obtained from the undermined edges of ulcerative lesions were soaked into 1 ml ethanol immediately after sampling and stored in polypropylene collecting tubes at -20°C, protected from light, until analysis. Ethanol solutions of exudates were concentrated to a volume of 500 μl then processed to TLC-Fluo and HPLC/MS/MS analyses.

Table 1. Characteristics of patients and controls.

| Ulcer exudates | Number | Age* | Gender (M/F)b | HPLCc |
|----------------|--------|------|---------------|-------|
| Before         | 6      | 23.3 | 0/6           | 50    |
| During         | 13     | 16.7 | 7/6           | 100   |
| After          | 4      | 20.7 | 0/4           | 100   |
| Ctrl           | NA     | NA   | NA            | NA    |

| Serum samples (Ghana) | Number | Age* | Gender (M/F)b | TLC-Boe |
|-----------------------|--------|------|---------------|---------|
| Before*               | 8      | 13.1 | 5/3           | 50      |
| During*               | 6      | 19.8 | 3/3           | 17      |
| After                 | 4      | 29.2 | 2/2           | 25      |
| Ctrl                  | 1      | 32.0 | 1/0           | 0       |

| Serum samples (Ivory Coast) | Number | Age* | Gender (M/F)b | HPLCc |
|-----------------------------|--------|------|---------------|-------|
| Before                      | 5      | ND   | ND            | 60    |
| During                      | 8      | 23.5 | 1/3           | 12    |
| After                       | 4      | 19.7 | 2/5           | 0     |
| Ctrl                        | 3      | ND   | ND            | 0     |

| Mononuclear cells (Ghana) | Number | Age* | Gender (M/F)b | HPLCc |
|---------------------------|--------|------|---------------|-------|
| Before*                   | 15     | 21.1 | 9/6           | 0     |
| During                     | 20     | 24.9 | 9/11          | 0     |
| After                      | 7      | 24.7 | 4/3           | 0     |
| Ctrl                       | 4      | 14.2 | 1/3           | 0     |

| Mononuclear cells (Ivory Coast) | Number | Age* | Gender (M/F)b | HPLCc |
|---------------------------------|--------|------|---------------|-------|
| Before                          | 3      | 33.3 | 1/2           | 0     |
| During                          | 4      | 26.0 | 2/2           | 0     |
| After                           | 5      | 19.6 | 0/5           | 0     |
| Ctrl                            | 4      | 30.0 | 2/2           | 0     |

*Mean age (year).
*M, males; F, females.
*Percentage of samples with signal superior to background.
*Newly diagnosed.
*Treated for 2 to 8 weeks.
*Completed the treatment.
*Healthy control from the same endemic zone.
NA: not applicable; ND: not determined.
doi:10.1371/journal.pntd.0001237.t001

Serum samples
Lipids were extracted from serum samples (1 ml) by sequential addition of 4/1 MeOH (v/v), 1/1 CHCl3 (v/v), and 3/1 H2O (v/v), each step being followed by thorough mixing. The upper aqueous phase was discarded and the bottom organic phase transferred to a glass tube containing 3 ml of MeOH. The resulting solution was centrifuged to sediment insoluble particulate matter. The soluble organic phase was then dried and the resulting product re-suspended in ethanol for TLC-Fluo and HPLC/MS/MS analyses.

Mononuclear cell pellets
Mononuclear cells were isolated from 10 ml whole blood by differential sedimentation on Ficoll-Hypaque (GE Healthcare). Cells pellets were then dried and stored at -20°C until lipid analysis. Total lipids were extracted from cell pellets by addition of 1 ml 2/1 CHCl3/MeOH (v/v) for 48 h at 4°C. The organic phase was recovered by
addition of 20% H$_2$O (w/v), dried and the resulting product resuspended in ethanol for TLC-Fluo and HPLC/MS/MS analyses.

Results

Efficacy of mycolactone extraction from biological samples

We first compared the TLC-Fluo and HPLC/MS/MS approaches for mycolactone detection in the 0–500 ng range, using purified mycolactone as a reference. Both methods were highly sensitive, yielding a detectable signal with only 10 ng mycolactone (Fig. 1A,B). However the fluorescent signal of mycolactone modified by coupling to 2-naphtalene boronic acid was not a linear function of its concentration, whereas the areas of mycolactone elution peaks in HPLC were proportional to mycolactone concentration in the 10–500 ng domain (Fig. 1C). To evaluate the efficacy of mycolactone extraction from biological samples, serum samples (1 ml) or mononuclear cell pellets (10$^6$ cells) were spiked with purified mycolactone. We then proceeded to solvent extraction as described in the Methods section, and estimated the proportion of recovered mycolactone using the above-described standard curves. The estimated yields of extraction with organic solvents were of 20% from cell pellets and 10% from serum samples (not shown). Several combinations of solvents were tried that did not improve the recovery of mycolactone.

Detection of mycolactone in ulcer exudates

Exudate samples were split into two equivalent parts, which were analyzed in parallel for the presence of mycolactone by HPLC/MS/MS or TLC-Fluo. We could not conclude on the presence of mycolactone by TLC-Fluo, because of the co-migration of auto-fluorescent compounds (Fig. 2A). Using HPLC determination, elution peaks corresponding to mycolactone were observed in 3/6 newly diagnosed patients and all patients undergoing or completing their course of antibiotic treatment (13/13, 4/4 respectively) (Table 1). To confirm that they effectively contained mycolactone, elution peaks were collected in three patients, and analyzed by MS/MS. In all of them, the characteristic spectrum of mycolactone parent ion (m/z 765) and products was observed [14], demonstrating the presence and structural integrity of mycolactone in ulcer exudates (Fig. 2B). Notably, the presence of mycolactone in these samples persisted during and after completion of antibiotic therapy (Fig. 3).

Detection of mycolactone in blood mononuclear cells

Since our previous experiments in experimentally infected mice demonstrated the presence of mycolactone in circulating mononuclear cells, we analyzed this biological material in BU patients. Whole blood (10 ml) was collected in patients from Ghana and Ivory Coast at various stages of the disease (Table 1) and mononuclear cells were isolated by Ficoll gradient centrifugation. However, no mycolactone could be identified in any of the samples tested (n = 52) by TLC-Fluo or HPLC/MS/MS (not shown).

Detection of mycolactone in serum

Total lipids were extracted from 1 ml serum samples then analyzed by TLC-Fluo (Ghana samples) or HPLC (Ivory Coast samples) (Table 1). Again, the identification of mycolactone by TLC-Fluo was difficult, because of the co-migration of auto-fluorescent compounds (Fig. 4A). Using HPLC, mycolactone detection was also hampered by the co-elution of UV-absorbing contaminants (Fig. 4B). We could nevertheless identify mycolactone-like peaks in 3/5 newly diagnosed patients, 1/8 patients undergoing treatment, and 0/4 patients completing treatment. These peaks were collected in two patients and analyzed by MS/MS. In both of them, the characteristic spectrum of mycolactone parent and product ions could be observed (Fig. 4B). The concentration of circulating mycolactone was evaluated in positive samples by measuring the area of mycolactone elution peaks and subtracting the mean signal of two healthy controls (Fig. 5). Calculated values were in the 40–200 ng/ml range.

Discussion

In the present study, we investigated whether mycolactone was detectable in easily accessible samples of BU patients. We used two chemical approaches, both requiring the extraction of total lipids by organic solvents. The efficacy of this extraction step, as measured by addition of pure mycolactone to control samples, was mediocre and reduced dramatically the sensitivity of the following
TLC-Fluo and HPLC determinations. An explanation for such a limited yield of extraction may be that mycolactone associates with biomolecules preventing solvent access, a possibility that we are currently testing by studying the impact of various thermal and enzymatic treatments. If the efficacy and selectivity of mycolactone extraction can be improved, the recently described and field-friendly TLC-Fluo detection method may still be an option for mycolactone-based point-of-care diagnostic tests. If not, it will be necessary to design alternative approaches that do not require this purification step.

Using HPLC/MS/MS, we could demonstrate that mycolactone gains access to the peripheral blood of human patients. In previous studies in the mouse model, we detected structurally intact mycolactone in mononuclear cell fractions of pooled blood samples harvested from mice subcutaneously injected with mycolactone, or experimentally infected with *M. ulcerans* [12]. In the present work, mycolactone could not be identified in blood mononuclear cells. This cell subpopulation was isolated from 10 ml whole blood, and we estimated the maximum yield of mycolactone extraction from mononuclear cell pellets to 20%. If mycolactone effectively reaches blood mononuclear cells in human patients, its cellular concentration may be too low to be detected in the accessible volume of blood. Alternatively, mycolactone may be unstable in the conditions used in the present study to isolate and store mononuclear cell pellets.

In contrast, we were able to demonstrate the presence of structurally intact mycolactone in the serum of 3/5 newly diagnosed BU patients (Table 1). This novel information provides the essential proof of concept for the design of BU diagnostic tests based on mycolactone detection in peripheral blood. Since mycolactone is extracted from serum samples with low efficacy, the number of positive samples and the calculated concentrations of circulating mycolactone are probably underestimated in this study. Whether mycolactone kinetics in serum could be employed to monitor the response to antibiotic treatment will certainly be interesting to investigate further. Our preliminary results suggest

---

**Figure 4. Structurally intact mycolactone is detected in serum samples of BU patients.** (A) Example of the fluorescence signals given by serum samples (lanes 2–9), following lipid extraction and analysis by TLC-Fluo. Controls include 1 μg pure mycolactone (lane 1), lipids extracted from a negative serum (lane 10), and then spiked with 1 μg mycolactone (lane 11). The band corresponding to mycolactone is designated by an asterisk. (B) Representative HPLC elution profiles of lipids extracted from serum samples are shown for one healthy control out of 5, and one BU patient among the 4 positive ones. The corresponding MS/MS spectra show the presence of the parent and product ions of mycolactone in this positive sample. Similar results were obtained in a second one.

doi:10.1371/journal.pntd.0001237.g004
that, in spite of a sustained presence in ulcer exudates (Fig. 3), mycolactone concentration (Fig. 5) showed a tendency to decrease in the serum during antibiotic therapy. If confirmed by longitudinal studies, the decay of circulating mycolactone during antibiotic therapy would provide an explanation to the recovery of cellular immune responses during treatment [15,16] and after surgical excision of BU lesions [17].

Here we considered patients at ulcerative stages of the disease. BU is usually diagnosed on the basis of clinical symptoms, as the identification of M. ulcerans by means of cultures or PCR requires dedicated facilities and specialized equipment (reviewed in [18]). Common differential diagnoses of BU include other tropical ulcers (venous, phagedenic, neurogenic), leishmaniasis, yaws and squamous cell carcinoma. The presence of biologically active mycolactone was recently demonstrated in skin biopsies of BU patients [11]. Here we show that mycolactone can be detected in ulcer exudates obtained non-invasively from wound swabs, in a structurally intact form and at concentrations in the 50–200 ng/ml range, which strongly suggests that mycolactone detection in exudates may be of interest for differential diagnosis of the ulcerative forms. Whether mycolactone is present at pre-ulcerative stages in fine-needle aspirates and in the peripheral blood is currently under investigation.

We observed significant amounts of mycolactone in exudates at the end of antibiotic therapy. In some instances, antibiotic pressure may not efficiently block, and could even enhance the production of a toxin. For instance, in patients infected with E. coli O157:H7, the use of antimicrobials has been discouraged because it stimulates toxin production and augments the risk of detrimental, or even fatal complications [19]. Whether antibiotics promote the expression of mycolactone by M. ulcerans is unknown. However, their efficacy at killing M. ulcerans is not questionable with 4 weeks of treatment leading to culture negativity [3,4,5,6,7]. Although a stimulatory effect of antibiotics on mycolactone production cannot be excluded, our observation suggests that mycolactone persists in cutaneous tissues after the demise of M. ulcerans. Since mycolactone displays inherent ulcerative properties [8], this phenomenon may explain why some BU take considerable time to heal despite remaining culture negative.

The administration of antibiotics is sometimes associated with adverse reactions, excessive inflammation and pain sensation, that paradoxically make the symptoms of infection worse [20]. It is possible that in BU, like in the Jarish-Heckheimer reaction in syphilis or in relapsing fevers, efficient killing of M. ulcerans by antibiotics leads to the sudden and massive release of bacterial antigens locally that act as immuno-stimulants. Mycolactone displays potent immunosuppressive properties in vitro that are thought to contribute to the cellular response defects of BU patients [21,22,23,24,25]. The rapid decline of mycolactone from the systemic circulation during treatment may thus provoke exuberant inflammatory responses at the level of the lesions, and cause the paradoxical reactions observed in BU disease [26].

**Author Contributions**

Conceived and designed the experiments: FSS ROP MD LGM RVC CD. Performed the experiments: FSS FLC NA IGB PL YA. Analyzed the data: FSS FLC NA CD. Contributed reagents/materials/analysis: MW-J. Wrote the paper: CD.

---

**References**

1. Silva MT, Portaels F, Pedrosa J (2009) Pathogenetic mechanisms of the intracellular parasite Mycobacterium ulcerans leading to Buruli ulcer. Lancet Infect Dis 9: 699–710.
2. Walsh DS, Portaels F, Meyers WM (2010) Recent advances in leprosy and Buruli ulcer (Mycobacterium ulcerans infection).Curr Opin Infect Dis 23: 445–455.
3. Etuaful S, Carbonnelle B, Grosset J, Lucas S, Horsfield C, et al. (2005) Efficacy of the combination rifampin-streptomycin in preventing growth of Mycobacterium ulcerans. Antimicrob Agents Chemother 51: 3678–3685.
4. Chaury A, Ardant MF, Adeye A,埃文特 H, Guedenon A, et al. (2007) Promising clinical efficacy of streptomycin-rifampin combination for treatment of Buruli ulcer (Mycobacterium ulcerans disease). Antimicrob Agents Chemother 51: 4029–4035.
5. Johnson PD (2010) Should antibiotics be given for Buruli ulcer? Lancet 375: 618–619.
6. Sarfo FS, Phillips R, Asiedu K, Ampadu E, Bobi N, et al. (2010) Clinical efficacy of combination rifampin and streptomycin for treatment of Mycobacterium ulcerans disease. Antimicrob Agents Chemother 54: 3678–3685.
7. Nieuwhuis WA, Sienstra Y, Thompson WA, Awoah PC, Abas KM, et al. (2010) Antimicrobial treatment for early, limited Mycobacterium ulcerans infection: a randomised controlled trial. Lancet 375: 664–672.
8. George KM, Chatterjee D, Gunawardana G, Welty D, Hayman J, et al. (1999) Mycolactone: a polyketide toxin from Mycobacterium ulcerans required for virulence. Science 283: 854–857.
9. Hong H, Demail C, Pilet SJ, Leadlay PF, Sinear T (2008) Mycolactones: immunosuppressive and cytotoxic polyketides produced by aquatic mycobacteria. Nat Prod Rep 25: 447–454.

10. Demail C, Sinear TP, Cole ST (2009) Buruli ulcer: reductive evolution enhances pathogenicity of Mycobacterium ulcerans. Nat Rev Microbiol 7: 50–60.

11. Sarfo FS, Phillips RO, Rangers B, Mahrous EA, Lee RE, et al. (2010) Detection of Mycolactone A/B in Mycobacterium ulcerans-infected human tissue. PLoS Negl Trop Dis 4: e577.

12. Hong H, Coutanceau E, Leclerc M, Caleechurn L, Leadlay PF, et al. (2008) Mycolactone diffuses from Mycobacterium ulcerans-infected tissues and targets mononuclear cells in peripheral blood and lymphoid organs. PLoS Negl Trop Dis 2: e325.

13. Spangenberg T, Kiabi Y (2010) Highly sensitive, operationally simple, cost/time effective detection of the mycolactones from the human pathogen Mycobacterium ulcerans. Chem Commun 46: 1410–1412.

14. Hong H, Sinear T, Shelon P, Spencer JB, Leadlay PF (2003) Structure elucidation of a novel family of mycolactone toxins from the frog pathogen Mycobacterium sp. MU129FXT by mass spectrometry. Chem Commun 14: 4306–4308.

15. Phillips R, Sarfo FS, Guerin-Mace L, Decalf J, Wambrough-Jones M, et al. (2009) Immunosuppressive signature of cutaneous Mycobacterium ulcerans infection in the peripheral blood of patients with Buruli ulcer disease. J Infect Dis 200: 1673–84.

16. Sarfo FS, Phillips RO, Ampadu E, Sarpong F, Adentew E, et al. (2009) Dynamics of the cytokine response to Mycobacterium ulcerans during antibiotic treatment for M. ulcerans disease (Buruli ulcer) in humans. Clin Vaccine Immunol 16: 61–65.

17. Yeboah-Manu D, Peuhu E, Mensah-Quainoo E, Azande-Pok A, Ofosu-Adjei D, et al. (2006) Systemic suppression of interferon-gamma responses in Buruli ulcer patients resolves after surgical excision of the lesions caused by the extracelular pathogen Mycobacterium ulcerans. J Leukoc Biol 79: 1150–1156.

18. Beissner M, Herbinger KH, Bretzel G (2010) Laboratory diagnosis of Buruli ulcer disease. Future Microbiol 5: 363–370.

19. Callacry TR, Anderson RC, Edrington TS, Jung YS, Bischoff KM, et al. (2004) Effects of sodium chloride on toxin production by Escherichia coli O157:H7. Curr Issues Intest Microbiol 5: 19–22.

20. Gordon CL, Bantine JA, Hayman JA, Lavender CJ, Fyfe JA, et al. (2010) Alloral antibiotic treatment for buruli ulcer: a report of four patients. PLoS Negl Trop Dis 4: e770.

21. Pahlevan AA, Wright DJ, Andrews C, George KM, Small PL, et al. (1999) The inhibitory action of Mycobacterium ulcerans soluble factor on monocyte/T cell cytokine production and NF-kappa B function. J Immunol 163: 3928–3935.

22. Boulkroun S, Guerin-Mace L, Thoulouze MI, Monot M, Merckx A, et al. (2010) Mycolactone suppresses T cell responsiveness by altering both early signaling and posttranslational events. J Immunol 184: 1436–1443.

23. Coutanceau E, Decalf J, Martino A, Babon A, Winter N, et al. (2007) Selective suppression of dendritic cell functions by Mycobacterium ulcerans toxin mycolactone. J Exp Med 204: 1395–1403.

24. Coutanceau E, Marsollier L, Brosch R, Perret E, Gosrens P, et al. (2005) Modulation of the host immune response by a transient intracellular stage of Mycobacterium ulcerans: the contribution of endogenous mycolactone toxin. Cell Microbiol 7: 1187–1196.

25. Simmonds RE, Lali FV, Smallie T, Small PL, Foxwell BM (2009) Mycolactone inhibits monocyte cytokine production by a posttranscriptional mechanism. J Immunol 182: 2194–2202.

26. Chute D, Pluschke G (2009) Immunosuppression and treatment-associated inflammatory response in patients with Mycobacterium ulcerans infection (Buruli ulcer). Expert Opin Biol Ther 9: 187–200.