The Synthesis of L-Alanyl and β-Alanyl Derivatives of 2-Aminoacridone and Their Application in the Detection of Clinically-Important Microorganisms

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Abstract

In clinical microbiology the speed with which pathogenic microorganisms may be detected has a direct impact on patient health. One important strategy used in the laboratory is the growth of cultures in the presence of an enzymatic substrate which, once transformed by the appropriate microbial enzyme, generates a detectable colour or fluorescence output. Such substrates have previously been prepared by our group and others and are available as commercial diagnostic kits, however they all suffer from some degree of diffusion when used in a solid growth medium. This diffusion complicates the detection and differentiation of species in polymicrobial cultures and so we sought to improve on our previous work. In this work we have prepared and evaluated a series of novel fluorogenic enzyme substrates based on N-substituted-2-aminoacridones. All of the prepared substrates were found to be suitable for the detection and differentiation of certain microorganisms, however those based on the 2-amino-10-benzylacridone core in particular showed no apparent diffusion when incorporated into solid growth media. On transformation these substrates generated brightly fluorescent colonies that are clearly contrasted with the background medium due to the difference in emission wavelength (λem 445–450 nm for the substrate, λem 550 nm for the product). Here we have shown that our L-alanyl aminopeptidase substrate, 2-(N-L-alanyl)-10-benzylacridone, is particularly suited to the detection of Gram-negative bacteria, and our β-alanyl aminopeptidase substrate, 2-(N-β-alanylmino)-10-benzylacridone, to the detection of Pseudomonas aeruginosa and Serratia marcescens when grown on solid media incorporating these substrates. The resulting fluorophore shows no apparent diffusion from the colonies of interest, and the enhanced sensitivity offered by fluorescent emission may allow for the detection of these organisms as microcolonies using automated fluorescence microscopy.
section. bioMerieux staff also provided direction on study design and performed some analysis using the synthetic substrates described in this manuscript. Northumbria University provided facilities for synthesis and analysis (performed by GT under the direction of SPS and ALJ). Freeman Hospital staff (JDP) contributed to the study design and evaluation of the substrates.

**Competing Interests:** Two of the authors (MC and SO) are employed by bioMerieux, a co-funder in this work. Two of the authors (SPS and GT) and employed by Northumbria University, a co-funder of this work. One of the authors (JDP) is employed by the Freeman Hospital, and receives funding from bioMerieux. This does not alter the authors' adherence to PLOS ONE policies on sharing data and materials. There are no patents, products in development, or marketed products to declare.

**Introduction**

The use of enzymatic substrates as tools for the detection and identification of clinically-important microorganisms is a subject of particular interest in the health-care sector [1–3]. In diagnostic microbiology, the detection of specific types of aminopeptidase activities has proved useful. Thus, Gram-positive and Gram-negative microorganisms can often be differentiated by their L-alanine aminopeptidase activities [4,5]. This enzyme is ubiquitous in Gram-negative microorganisms whereas it is usually absent from most Gram-positive microorganisms. β-Alanly aminopeptidase has been detected in *Pseudomonas aeruginosa*, a common respiratory pathogen in cystic fibrosis patients [6].

One important strategy that has been utilised for microorganism detection and identification is to grow the microorganisms in the presence of an enzymatic substrate; microorganisms which possess the appropriate enzymes can then transform the substrate into a detectable product. The release of highly fluorescent heterocyclic amines from weakly fluorescent enzyme substrates has been employed as a protocol for detecting aminopeptidase activities, as illustrated in Fig 1.

Substrates of 7-amino-4-methylcoumarin are currently used in commercial diagnostic kits, and we have previously described the synthesis of fluorogenic L-alanyl and β-alanyl aminopeptidase substrates derived from 2-(2-aminophenyl)benzothiazoles and 2-(2-aminophenyl)benzoxazoles and their evaluation against a range of clinically-important microorganisms (Fig 2) [7–9]. Each of these substrates generates a fluorescent product which is yellow in colour (reducing the impact of cellular autofluorescence) and which does not appear to diffuse (providing high contrast to the growth medium).

**Methods**

The required core amines were readily prepared by alkylation of 2-nitroacridone [10] under basic conditions followed by reduction of the nitro-group (see ESI for synthetic procedures and characterisation data). The 2-aminoacridones were converted into their corresponding Boc-protected amino acid derivatives as depicted in Fig 3. Deprotection of these Boc-protected compounds using trifluoroacetic acid afforded the desired aminopeptidase substrates and .

The fluorescence excitation and emission wavelengths for the core amines and the substrates, 7a-c, 9 and 11 in ethanol are collected in Table 1. Solutions of the core amines and the substrates, 7a-c, 9 and 11 in ethanol are collected in Table 1. Solutions of the core amines all
displayed yellow fluorescence ($\lambda_{em}$ 545–546 nm). Conjugation through the 2-amino group significantly reduces the Stokes’ shift of the emission, and solutions of the substrates 7a-c, 9 and 11 showed blue fluorescence ($\lambda_{em}$ 445–453 nm). Fig 4 depicts ethanolic solutions of amine 5c and its corresponding L-alanyl aminopeptidase substrate 7c when viewed under UV illumination at 365 nm.

**Microbiological Evaluation and Discussion**

The substrates 7a-c, 9 and 11 were evaluated in Columbia agar medium against a panel of clinically-important microorganisms including 10 Gram-negative bacteria, 8 Gram-positive bacteria and 2 yeasts (see ESI for microbiological procedures). The growth of the microorganisms after incubation (18 h, 37°C) was compared to a control in which no substrate was present. Table 2 depicts the extent of growth of the various Gram-negative microorganisms and the fluorescence generated by hydrolysis of the substrates by L-alanyl and $\beta$-alanyl aminopeptidases (full results are supplied in the ESI). The substrates were all evaluated at a concentration of 50 mg L$^{-1}$ and the N-benzylated, L-alanyl substrate 7c was also evaluated at a concentration of 100 mg L$^{-1}$.

In the control, Gram-negative microorganisms exhibited strong growth (all graded ++) whereas Gram-positive microorganisms and yeasts only displayed moderate growth (all graded +). The majority of Gram-negative microorganisms exhibited strong growth in the presence of the substrates when the substrate concentration was 50 mg L$^{-1}$, with the exception of *Escherichia coli* and *Klebsiella pneumoniae* which did not grow in the presence of substrates 7a and 7b. The growth of the Gram-positive microorganisms was frequently diminished in comparison with the control, indicating an inhibitory effect of some of the substrates on some of these
bacteria. There was a clear inhibitory effect on all of the microorganisms by substrate 7c at a concentration of 100 mg L\(^{-1}\) in comparison with the growth observed at a concentration of 50 mg L\(^{-1}\).

All of the L-alanyl aminopeptidase substrates were able to produce bright yellow fluorescent colonies with most Gram-negative bacteria when viewed under UV light (365 nm), indicating hydrolysis of the substrate had occurred. The background fluorescence was blue as a consequence of the blue fluorescence of the unreacted substrate but there was a clear contrast between the yellow and blue fluorescence of the colonies and the background respectively. This is illustrated in Fig 5. Enzyme substrates for microorganism detection are generally designed to be weakly fluorescent or non-fluorescent, only producing strong fluorescence after release of...

Fig 3. Synthesis of aminopeptidase substrates. Reagents and conditions: (i) N-Boc-protected amino acid, N-methyl morpholine, isobutyl chloroformate, anhydrous THF, -12°C to RT overnight; (ii) TFA, RT, 2 h.

doi:10.1371/journal.pone.0158378.g003

Table 1. Excitation and emission wavelengths of amines 5a-c and substrates derived from these amines in ethanol solution (all 1 mg L\(^{-1}\)).

| Amine | \(\lambda_{ex}\) | \(\lambda_{em}\) | Substrate | \(\lambda_{ex}\) | \(\lambda_{em}\) |
|-------|----------------|----------------|-----------|----------------|----------------|
| 5a    | 434           | 546           | 7a        | 418           | 453           |
| 5b    | 435           | 545           | 7b        | 401,418       | 445           |
| 5c    | 429           | 550           | 7c        | 415           | 445           |
|       |               |               | 9         | 414           | 450           |
|       |               |               | 11        | 414           | 449           |

\(^a\)lit., 438 nm (EtOH).

doi:10.1371/journal.pone.0158378.t001
the core fluorophore, however the fluorescence of both substrate and product in this work is not a disadvantage because of their different emission wavelengths.

Substrate 7a gave strongly fluorescent yellow colonies with most Gram-negative microorganisms. Neither the Gram-positive microorganisms nor the yeasts produced fluorescent colonies with this substrate. The activity of substrate 7b was broadly similar to substrate 7a, but the fluorescence observed with the Gram-negative microorganisms was generally weaker. Blue fluorescent colonies were seen with *Burkholderia cepacia* and this was attributed to concentration of the substrate within the colonies without hydrolysis occurring. Substrate 7c gave fluorescent yellow colonies of variable intensities with the majority of the Gram-negative microorganisms, and weak, yellow fluorescent colonies with two of the Gram-positive microorganisms. At a higher substrate concentration (100 mg L⁻¹), this substrate was more inhibitory to microorganism growth and hence was observed to be more selective with six of the ten Gram-negative producing weakly fluorescent colonies. Good microorganism growth was seen with the di-L-alanyl substrate 9 with the majority of the Gram-negative microorganisms producing yellow fluorescent colonies of variable intensities.

As anticipated, the β-alanyl aminopeptidase substrate 11 gave intense, yellow fluorescent colonies with *Pseudomonas aeruginosa*. This substrate was also effective at detecting *Serratia marcescens*, a microorganism that is also known to possess β-alanyl aminopeptidase activity [4].
Each of these substrates are potentially useful for microbiological applications as they are cleaved to generate brightly fluorescent products that are clearly contrasted with the background culture medium due to their differing fluorescence emission wavelengths. Moreover, the fluorescent product released by hydrolysis remains restricted to the bacterial colonies and does not appear to diffuse through the agar. This lack of diffusion means that a colony expressing a specific aminopeptidase may be differentiated from a neighboring colony that does not express this activity using these substrates, such as in polymicrobial cultures routinely generated from food and clinical samples. These combined attributes make these substrates a useful alternative to many conventional substrates such as those based on 7-amino-4-methylcoumarin, which emits light in the blue region of the spectrum [11,12] and can suffer from a reduction in sensitivity by cellular autofluorescence.

Chromogenic culture media are widely used for detection of bacterial pathogens as they offer high specificity by targeting specific enzyme activities using chromogenic enzymes.

### Table 2. Growth and fluorescence of Gram-negative microorganisms in the presence of substrates 7a-c.

| Microorganism / Reference | Control | Substrate 7a | Substrate 7b | Substrate 7c |
|---------------------------|---------|--------------|--------------|--------------|
|                           | Growthb | Fluorescenceb | Growthb | Fluorescenceb | Growthb | Fluorescenceb |
| **Microorganism** / **Reference** |  |  |  |  |  |  |
| 1. *Escherichia coli* NCTC 10418 | ++ | - | - | - | ++ | ++ yellow |
| 2. *Klebsiella pneumoniae* NCTC 9528 | ++ | - | - | - | ++ | - |
| 3. *Providencia rettgeri* NCTC 7475 | ++ | ++ | ++ yellow | ++ | + yellow | ++ | ++ yellow |
| 4. *Enterobacter cloacae* NCTC 11936 | ++ | ++ | ++ yellow | ++ | + yellow | ++ | ++ yellow |
| 5. *Serratia marcescens* NCTC 10211 | ++ | ++ | ++ yellow | ++ | + yellow | ++ | ++ yellow |
| 6. *Salmonella typhimurium* NCTC 74 | ++ | ++ | ++ yellow | ++ | + yellow | ++ | +/- yellow |
| 7. *Pseudomonas aeruginosa* NCTC 10662 | ++ | ++ | ++ yellow | ++ | + yellow | ++ | ++ yellow |
| 8. *Yersinia enterocolitica* NCTC 11176 | ++ | ++ | + yellow | ++ | + yellow | ++ | + yellow |
| 9. *Burkholderia cepacia* NCTC 10743 | ++ | ++ | ++ yellow | ++ | + blue | ++ | ++ blue |
| 10. *Acinetobacter baumannii* NCTC 12156 | ++ | ++ | ++ yellow | - | - | + | + yellow |

**Background**

- ++ strong growth, + moderate growth, — no growth.
- ++ strong fluorescence, + moderate fluorescence, +/- weak fluorescence, — no fluorescence.

*a* NCTC: National Collection of Type Cultures; ATCC: American Type Culture Collection; NCPF: National Collection of Pathogenic Fungi.

| doi:10.1371/journal.pone.0158378.t002 |

**Fig 5. Colonies of*S. aureus* (left) and *E. coli* (right) after incubation in the presence of substrate 7c.**

doi:10.1371/journal.pone.0158378.g005
substrates. (e.g. ref. [13]). For example, the β-alanyl aminopeptidase activity of *Pseudomonas aeruginosa* has been targeted in a chromogenic culture medium for detection of this important pathogen [14]. Fluorescent substrates such as 11 may allow a similar approach with the generation of a fluorescent (rather than a chromogenic) product, and these are currently not widely used in solid media. The enhanced sensitivity offered by fluorescence and high-contrast afforded by these substrates may allow for the specific detection of pathogenic bacteria with reduced time-to-detection using automated detection of fluorescent microcolonies before they are visible to the naked eye [15].

**Supporting Information**

S1 File. Supporting information. (DOCX)

**Acknowledgments**

We thank the EPSRC Mass Spectrometry Centre, Swansea, for high-resolution mass spectra.

**Author Contributions**

Conceived and designed the experiments: MC ALJ SO JDP GT SPS. Performed the experiments: MC JDP GT. Analyzed the data: MC ALJ SO JDP GT SPS. Contributed reagents/materials/analysis tools: MC JDP GT SPS. Wrote the paper: JDP GT SPS.

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