A Metabolomics and Molecular Networking Approach to Elucidate the Structures of Secondary Metabolites Produced by Serratia marcescens Strains

Tanya Clements 1, Marina Rautenbach 2, Thando Ndlovu 1, Sehaam Khan 3 and Wesaal Khan 1*

1Department of Microbiology, Faculty of Science, Stellenbosch University, Stellenbosch, South Africa, 2BioPep Peptide Group, Department of Biochemistry, Faculty of Science, Stellenbosch University, Stellenbosch, South Africa, 3Faculty of Health Sciences, University of Johannesburg, Doornfontein, South Africa

An integrated approach that combines reverse-phase high-performance liquid chromatography (RP-HPLC), electrospray ionization mass spectrometry, untargeted ultra-performance liquid chromatography coupled to tandem mass spectrometry (UPLC-MSE) and molecular networking (using the Global Natural Products Social molecular network platform) was used to elucidate the metabolic profiles and chemical structures of the secondary metabolites produced by pigmented (P1) and non-pigmented (NP1) Serratia marcescens (S. marcescens) strains. Tandem mass spectrometry-based molecular networking guided the structural elucidation of 18 compounds for the P1 strain (including 6 serratamolides, 10 glucosamine derivatives, prodigiosin and serratiochelin A) and 15 compounds for the NP1 strain (including 8 serratamolides, 6 glucosamine derivatives and serratiochelin A) using the MSE fragmentation profiles. The serratamolide homologues were comprised of a peptide moiety of two L-serine residues (cyclic or open-ring) linked to two fatty acid chains (lengths of C10, C12, or C12:1). Moreover, the putative structure of a novel open-ring serratamolide homologue was described. The glucosamine derivative homologues (i.e., N-butylglucosamine ester derivatives) consisted of four residues, including glucose/hexose, valine, a fatty acid chain (lengths of C13 – C17 and varying from saturated to unsaturated) and butyric acid. The putative structures of seven novel glucosamine derivative homologues and one glucosamine derivative congener (containing an oxo-hexanoic acid residue instead of a butyric acid residue) were described. Moreover, seven fractions collected during RP-HPLC, with major molecular ions corresponding to prodigiosin, serratamolides (A, B, and C), and glucosamine derivatives (A, C, and E), displayed antimicrobial activity against a clinical Enterococcus faecalis S1 strain using the disc diffusion assay. The minimum inhibitory and bactericidal concentration assays however, revealed that prodigiosin exhibited the greatest antimicrobial potency, followed by glucosamine derivative A and then the serratamolides (A, B, and C). These results provide crucial insight into the secondary metabolic profiles of pigmented and non-pigmented S. marcescens strains.
and confirms that *S. marcescens* strains are a promising natural source of novel antimicrobial metabolites.

**Keywords:** pigmented and non-pigmented *Serratia marcescens*, molecular networking, prodigiosin, serratamolide, glucosamine derivative, UPLC-MS$^E$

**INTRODUCTION**

*Serratia* species are facultatively anaerobic, Gram-negative rods that are classified into the Enterobacteriaceae family (Grimont and Grimont, 2015). The type strain of this genus, *Serratia marcescens* (*S. marcescens*), is widely known to produce a distinctive red pigment referred to as prodigiosin (Su et al., 2016). *Serratia plymuthica* (*S. plymuthica*), *Serratia rubidaea* (*S. rubidaea*), and *Serratia nematodiphila* (*S. nematodiphila*) have also been reported to synthesize this pigment, while non-pigmented strains of these species have been identified (Su et al., 2016). In recent years, pigmented and non-pigmented *Serratia* species have been recognized as a potential source of novel and structurally diverse bioactive secondary metabolites (Soenens and Imperial, 2019). These bioactive metabolites include the pigment prodigiosin as well as biosurfactants (lipopeptides and glycolipids), glucosamine derivatives, oocydin A, siderophores (such as serratiochelin A; Figure 1A), bacteriocins, carabapenem, althiomycin and serratin, among others (Clements et al., 2019a; Khilyas et al., 2019). Prodigiosin and serrawettins are two of the more extensively studied secondary metabolites due to their diverse biological activity and application as antitumor, antibacterial and antifungal agents (Stankovic et al., 2014; Eckelmann et al., 2018; Clements et al., 2019a).

Prodigiosin is a red, tripyrrole pigment (family of prodiginines) that is produced by several bacterial genera, such as *Serratia, Hahella, Streptomyces, Zooshikella, Vibrio,* and *Pseudomonas,* among others (Darshan and Manonmani, 2015). Prodigiosin has a general structure of 2-methyl-3-pentyl-6-methoxyprodiginine (Figure 1B) and is composed of three rings referred to as pyrrolic ring A, B, and C (Hage-Hülsmann et al., 2018; Yip et al., 2019). Research has indicated that it is synthesized via a bifurcated pathway encoded in a pigment (pig) gene cluster comprised of 14 genes (*pigABCDEFGHIJKLMN*) (Williamson et al., 2006). In contrast, serrawettins are non-ribosomally synthesized cyclodepsipeptides solely produced by several members of the *Serratia* genus. Serrawettins are classified as lipopeptides as they are composed of a hydrophilic peptide moiety and a hydrophobic fatty acid moiety. Three unique structures of serrawettins were initially identified, including serrawettin W1 (also referred to as serratamolide A), serrawettin W2 and serrawettin W3 (Wasserman et al., 1961; Matsuyama et al., 1985; Matsuyama et al., 1986; Matsuyama et al., 1992), while various analogues and homologues of serrawettin W1 and W2 have since been described (Dwivedi et al., 2008; Motley et al., 2016; Su et al., 2016). *Serratamolide A* and homologues of this lipopeptide (including serramolide B to G) are comprised of a cyclic peptide moiety of two L-serine amino acids linked to two β-hydroxy fatty acid moieties that may vary in length (C$_6$ to C$_{14}$) of the fatty acyl chain and may be saturated or unsaturated (Figure 1C; Dwivedi et al., 2008). In comparison, serrawettin W2 and W3 are comprised of five amino acids linked to one β-hydroxy fatty acid (Matsuyama et al., 1986; Matsuyama et al., 1992).

**FIGURE 1 |** General structures of (A) serratiochelin A, (B) prodigiosin, (C) serratamolide and (D) glucosamine derivative (adapted from Dwivedi et al., 2008; Lee et al., 2011; Seyedsayamdost et al., 2012). The $R_1$ and $R_2$ groups on the serratamolides and glucosamine derivative structures indicate the fatty acyl moieties and these chains may vary in length and saturation (saturated, mono-unsaturated or poly-unsaturated).
Glucosamine derivatives (also referred to as N-butyglycosamine ester derivatives) are non-ribosomally synthesized diaxylated peptidoglycosamine derivatives (Dviwedi et al., 2008; Khilyas et al., 2019) produced by *Serratia* species. They have not been extensively studied and to date only Dviwedi et al. (2008) characterized glucosamine derivatives produced by a pigmented *S. marcescens* SHHRE645 strain. The identified glucosamine derivative homologues (A to C) were comprised of four residues, including glucose, valine, butyric acid and a saturated or unsaturated β-fatty acid residue of varying length (Figure 1D). It was observed that the glucosamine derivatives were co-produced with serratamolide homologues. Although extensive research into the biological properties of glucosamine derivative homologues (A to C) is still required, Dviwedi et al. (2008) reported on the anti-mycobacterial activity for each homologue.

To date, several analytical methods have been utilized to elucidate the detailed structures and composition of secondary metabolites produced by *Serratia* species, including mass spectrometry (MS), liquid chromatography (LC), gas chromatography (GC), and nuclear magnetic resonance (NMR) (Dwivedi et al., 2008; Thies et al., 2014; Eckelmann et al., 2018). Chromatography methods are commonly coupled with MS for the reliable separation and identification of bioactive compounds (Fu et al., 2019). Moreover, tandem MS analysis allows for the detection and structural characterization of a broad range of compounds and distinguishes between closely related forms of the same compound produced by a bacterial strain (Knolhoff et al., 2015). For example, a study by Eckelmann et al. (2018) unraveled the secondary metabolic profile of a pigmented *S. marcescens* strain MSRBB2 using high performance LC (HPLC) linked to high resolution MS (HRMS) and HRMS mass spectrometry coupled to tandem mass spectrometry (UPLC-MS5) and molecular networking (using the GNPS molecular network platform) analysis. In addition, the susceptibility of a clinical *Enterococcus faecalis* (*E*. *faecalis*) strain to fractions collected during RP-HPLC analysis was evaluated using disc diffusion assays (all fractions) and broth microdilution assays to determine the minimum inhibitory concentration (MIC) and minimum bactericidal concentration (MBC) of selected fractions.

**MATERIALS AND METHODS**

**Bacterial Strains**

A previous study by Clements et al. (2019b) outlined the isolation of the P1 and NP1 *S. marcescens* strains from an oil refinery effluent sample and a river water sample, respectively as well as the molecular identification of the strains. The two *S. marcescens* strains were streaked from the glycerol stocks onto Nutrient agar (Merck, Johannesburg, South Africa) and were incubated at 30°C for 18–24 h. The *S. marcescens* P1 and NP1 strains were deposited in the South African Rhizobium Culture Collection (SARCC no. for P1: 3059; SARCC no. for NP1: 3060). The clinical *E. faecalis* S1 strain, used for the disc diffusion and MIC assays, was streaked from the glycerol stock onto tryptone soy agar (Biolab, Merck, Johannesburg, South Africa) supplemented with 6 g/L yeast extract (Biolab, Merck, Johannesburg, South Africa) (TSAYE0.6%) and was incubated at 37°C for 18–24 h. The clinical *E. faecalis* S1 strain is curated and accessible in the Water Resource Laboratory culture collection in the Department of Microbiology at Stellenbosch University.

**Production and Extraction of Secondary Metabolites**

The production and partial purification of secondary metabolites was performed as described by Clements et al. (2019b), with slight modifications to the scale of production (increased volume) and the use of peptone powder in culture media (Biolab, Merck, Johannesburg, South Africa). Briefly, seed cultures were grown overnight by inoculating the *S. marcescens* P1 and NP1 strains into 10 ml Luria Bertani broth (Biolab, Merck, South Africa) in triplicate and incubating on a test tube rotator (MRCLAB, London, United Kingdom) at 30°C. Each seed culture was subsequently inoculated into a 2 L baffled flask containing 500 ml Peptone Glycerol (PG, pH 7.2 ± 0.2) broth [composed of 5 g peptone powder and 10 ml glycerol (Promega, Wisconsin, United States) in 1 L distilled water] in triplicate, which was incubated on an orbital shaker (MRCLAB, London, United Kingdom) at 30°C for 120 h at 120 rpm. Following a 5 days incubation period, the P1 and NP1 broth cultures were centrifuged at 10,000 rpm for 20 min at 4°C to obtain the cell free supernatant. The cell free supernatants were lyophilized and dissolved in 70% HPLC-grade acetonitrile (Romil, Darmstadt, Germany) in analytical quality water (milliQ water) (v/v). The acetonitrile soluble fraction was transferred into a sterile McCartney bottle and lyophilized. This step was repeated...
Purification and Detection of Secondary Metabolites

The P1 and NP1 crude extracts were subjected to RP-HPLC (Finnigan Survey UV-VIS Plus detector, Thermo-Scientific, Waltham, MA, United States) analysis, in the Department of Microbiology, in order to obtain purified fractions. The lyophilized P1 and NP1 crude extracts were dissolved in 40% acetonitrile in milliQ water (v/v) to a concentration of 10.00 mg/ml and were injected into a Discovery BIO Wide Pore C18 HPLC column (10 μm, 250 × 10 mm; Sigma-Aldrich, St. Louis, United States). Liquid chromatography was conducted with milliQ water containing 0.1% trifluoroacetic acid (TFA; v/v; Sigma-Aldrich, St. Louis, United States) as solution A and acetonitrile containing 0.1% TFA (v/v) as solution B. An isocratic flow at 40% B from 0 to 2 min for sample loading was followed by a linear gradient from 40 to 95% B from 2 to 36 min. The column was washed for 2 min with 95% B and then regenerated with a reversed gradient from 95 to 40% B from 38 to 45 min. Chromatography of compounds was followed by continuous monitoring of the absorbance at 230 and 254 nm.

The absorbing peak fractions collected from the RP-HPLC were analyzed using a Waters Synapt G2 high resolution mass spectrometer fitted with a Z spray electrospray ionization source (Waters Corporation, Milford, United States) at the LCMS unit at the Central Analytical Facility (CAF, Stellenbosch University). For direct mass analysis, 3 μl of the sample [250 μg/ml; dissolved in 50% acetonitrile in milliQ water (v/v)] was injected into the ionization source at a flow rate of 0.3 ml/min. The analytes were subjected to a capillary voltage of 2.5 kV, cone voltage of 15 V, a source temperature of 120°C, desolvation gas in milliQ water (v/v) to a concentration of 250 μg/ml and were subjected to the Waters Synapt G2 high resolution mass spectrometer linked to an Acquty UPLC™ for UPLC-MS analysis. Three microlitres of each fraction (1.00 mg/ml) was separated on an UPLC C18 reverse-phase analytical column (Acquity UPLC® HSS T3, 1.8 μm particle size, 2.1 × 150 mm, Waters Corporation, Dublin, Ireland). The compound separation via UPLC was facilitated with milliQ water containing 0.1% (v/v) formic acid as solution A and acetonitrile containing 0.1% (v/v) formic acid as solution B. The gradient was developed at a flow rate of 0.300 ml/min as follows: 60% A from 0 to 0.5 min for loading, linear gradient from 20 to 80% (B) from 0.5 to 14 min and 0–100% (B) from 14 to 15 min. The ESI-MS instrument settings for monitoring the chromatography were as described for the direct mass analysis.

High resolution collisionally induced dissociation (CID) analysis was conducted in the MS\(^\text{E}\) mode (MS/MS) during the UPLC-MS analysis and was monitored on a second MS channel. The CID was conducted at a collision energy gradient of 20–70 eV at 1 s MS/MS scan time. Data was collected in the second mass analyser (MS2) through m/z range of 40–1,500 in centroid mode. The rest of the instrument settings were as described above. The UPLC-MS\(^\text{E}\) data was processed using Masslynx software version 4.1 (Waters Corporation).

Molecular Networking Analysis

The Waters RAW files for the P1 and NP1 fractions obtained after UPLC-MS\(^\text{E}\) analysis were converted into an Analysis Base File (ABF) format using Reiflycs Analysis Base File Converter before data processing. Ion chromatogram extraction, alignment and peak deconvolution of the ABF converted files was then conducted using MS-DIAL software version 4.24. The aligned results were exported as a mascot generic format (mgf) file for P1 and NP1 (Tsugawa et al., 2015). Thereafter, the mgf files for the two strains were uploaded to the GNPS platform (http://gnps.ucsd.edu) and a molecular network was created using the workflow published by Wang et al. (2016). Briefly, the data was filtered by removing all MS/MS fragment ions within +/−17 Da of the precursor m/z. The MS/MS spectra were window filtered by selecting only the top six fragment ions in the +/−50 Da window throughout the spectrum. The following parameters were then used in the workflow: precursor ion mass tolerance was set to 0.03 Da, MS/MS fragment ion tolerance of 0.02 Da, cosine score above 0.6 and minimum matched peaks of ten. Further, edges between two nodes were kept in the network only if each of the nodes appeared in each other’s respective top 10 most similar nodes. The spectra in the network were then searched against GNPS’ spectral libraries. All matches retained between network spectra and library spectra were required to have a score above 0.7 and at least six matched peaks (Wang et al., 2016). The output of the molecular network was visualized using Cytoscape version 3.8.0. The nodes (compounds) originating from media and solvent controls (acetonitrile) were excluded.

Analysis Utilizing Untargeted Ultra-performance Liquid Chromatography Linked to Tandem Mass Spectrometry

In order to elucidate the structure of each of the detected compounds, all fractions were subjected to untargeted ultra-performance liquid chromatography coupled to tandem mass spectrometry (UPLC-MS\(^\text{E}\)) analysis at the LCMS unit at the Central Analytical Facility (CAF, Stellenbosch University). The RP-HPLC purified fractions obtained from the P1 and NP1 crude extracts were prepared in 70% acetonitrile in milliQ water (v/v) to a concentration of 250 μg/ml and were subjected to the Waters Synapt G2 high resolution mass spectrometer linked to an Acquty UPLC™ for UPLC-MS analysis. Three microlitres of each fraction (1.00 mg/ml) was separated on an UPLC C18 reverse-phase analytical column (Acquity UPLC® HSS T3, 1.8 μm particle size, 2.1 × 150 mm, Waters Corporation, Dublin, Ireland). The compound separation via UPLC was facilitated with milliQ water containing 0.1% (v/v) formic acid as solution A and acetonitrile containing 0.1% (v/v) formic acid as solution B. The gradient was developed at a flow rate of 0.300 ml/min as follows: 60% A from 0 to 0.5 min for loading, linear gradient from 20 to 80% (B) from 0.5 to 14 min and 0–100% (B) from 14 to 15 min. The ESI-MS instrument settings for monitoring the chromatography were as described for the direct mass analysis.

High resolution collisionally induced dissociation (CID) analysis was conducted in the MS\(^\text{E}\) mode (MS/MS) during the UPLC-MS analysis and was monitored on a second MS channel. The CID was conducted at a collision energy gradient of 20–70 eV at 1 s MS/MS scan time. Data was collected in the second mass analyser (MS2) through m/z range of 40–1,500 in centroid mode. The rest of the instrument settings were as described above. The UPLC-MS\(^\text{E}\) data was processed using Masslynx software version 4.1 (Waters Corporation).

Molecular Networking Analysis

The Waters RAW files for the P1 and NP1 fractions obtained after UPLC-MS\(^\text{E}\) analysis were converted into an Analysis Base File (ABF) format using Reiflycs Analysis Base File Converter before data processing. Ion chromatogram extraction, alignment and peak deconvolution of the ABF converted files was then conducted using MS-DIAL software version 4.24. The aligned results were exported as a mascot generic format (mgf) file for P1 and NP1 (Tsugawa et al., 2015). Thereafter, the mgf files for the two strains were uploaded to the GNPS platform (http://gnps.ucsd.edu) and a molecular network was created using the workflow published by Wang et al. (2016). Briefly, the data was filtered by removing all MS/MS fragment ions within +/−17 Da of the precursor m/z. The MS/MS spectra were window filtered by selecting only the top six fragment ions in the +/−50 Da window throughout the spectrum. The following parameters were then used in the workflow: precursor ion mass tolerance was set to 0.03 Da, MS/MS fragment ion tolerance of 0.02 Da, cosine score above 0.6 and minimum matched peaks of ten. Further, edges between two nodes were kept in the network only if each of the nodes appeared in each other’s respective top 10 most similar nodes. The spectra in the network were then searched against GNPS’ spectral libraries. All matches retained between network spectra and library spectra were required to have a score above 0.7 and at least six matched peaks (Wang et al., 2016). The output of the molecular network was visualized using Cytoscape version 3.8.0. The nodes (compounds) originating from media and solvent controls (acetonitrile) were excluded.
from the original network in order to visualize the secondary metabolites derived from the P1 and NP1 strains.

**Antimicrobial Susceptibility Testing Disc Diffusion Assay**

The lyophilized fraction stocks (obtained after RP-HPLC) were dissolved in 100% dimethyl sulfoxide (DMSO; Sigma-Aldrich, St. Louis, United States) to a concentration of 20.00 mg/ml (stock concentration). Following solubility of each fraction, an aliquot of each fraction was diluted in milliQ water to a final concentration of 3.00 mg/ml (with a final DMSO concentration of 15%). These aliquots (at 3.00 mg/ml) were then subjected to antimicrobial testing using a standard disc diffusion assay as described by Das et al. (2008) and Ndlovu et al. (2017). Briefly, the *E. faecalis* S1 test strain was inoculated into 5 ml of tryptone soy broth (Biolab, Merck, Johannesburg, South Africa) supplemented with 6 g/L yeast extract (TSBYE0.6%) and was inoculated at 37°C for 18–24 h. Following incubation, 100 µl of the *E. faecalis* S1 suspension was spread plated onto TSBYE0.6% to create a microbial lawn. Thereafter, 6 mm filter paper discs (Oxoid, Basingstoke, United Kingdom) were placed onto the microbial lawn using a sterile needle and 50 µl of each fraction (3.00 mg/ml in 15% DMSO) was pipetted directly onto the filter paper (in triplicate) in order to create an antimicrobial disc. A negative control of 50 µl of 15% (v/v) DMSO was included and the plates were incubated for 24 h at 37°C.

**Minimum Inhibitory and Bactericidal Concentration**

Fractions were selected for further antimicrobial testing based on the results of the disc diffusion assay and the quantity of each fraction remaining. The fraction stocks were prepared in 100% DMSO (as outlined in 2.6.1) to 20.00 mg/ml and aliquots were prepared and diluted to 6.00 mg/ml (DMSO concentration of 30%) using milliQ water. A serial dilution of the fractions at 6.00 mg/ml was then conducted using 30% DMSO to obtain a concentration range of 1.50–60.00 mg/ml (DMSO concentration of 30%). The selected fractions (at various concentrations) were subjected to antimicrobial testing using a broth microdilution susceptibility assay as outlined by the European Committee on Antimicrobial Susceptibility Testing (European Committee on Antimicrobial Susceptibility Testing, 2018) and Magalhães and Nitschke et al. (2013). Briefly, *E. faecalis* S1 was inoculated into 5 ml of TSBYE0.6% and was incubated at 37°C for 18–24 h. Following overnight incubation, the culture was diluted to an optical density (OD) of approximately 0.08 at 625 nm corresponding to ~10^7 colony forming units (CFU)/ml. One hundred microlitres of each fraction (at the various concentrations) was dispensed into the respective wells of a Greiner CELLSTAR® 96-well culture plate (Merck, Johannesburg, South Africa) containing 100 µl of sterile TSBYE0.6% (resulting in a final test concentration of 0.75–3.00 mg/ml in 15% DMSO). All the wells of the microtiter plate were inoculated with 10 µl of the bacterial inoculum (final concentration of ~10^5 CFU/ml). Sterile broth, 15% DMSO and the OD adjusted inoculum were included in the assay as positive controls, while sterile broth with 15% DMSO was included as a negative control. All the tests were performed in triplicate and the microtiter plate was incubated for 18–24 h at 37°C. Following incubation, the absorbance was measured using a microtiter plate reader. The MIC was assigned as the lowest concentration of the respective compounds that reduced bacterial numbers by ≥90% (Yasir et al., 2019). The MBC was determined as previously described by Das et al. (2008). Briefly, after the MIC determination, the wells that showed ≥90% inhibition of growth were identified and 20 µl of each well was spot plated onto TSBYE0.6%. The lowest concentration showing no revival of the test culture was recorded as the MBC.

**RESULTS**

**Purification and Identification of Secondary Metabolites**

Reverse-phase HPLC was used to fractionate the secondary metabolites within the P1 and NP1 crude extracts. The RP-HPLC analysis of the P1 and NP1 crude extracts revealed *n* = 11 absorption peaks in the P1 crude extract and *n* = 8 absorption peaks in the NP1 crude extract, between 8 and 35 min. All the collected fractions were then subjected to ESI-MS analysis to detect potential metabolites produced by each strain. As a result, a combined total of 21 compounds, produced by the P1 and/or NP1 strains, were detected [numbered as compounds 1–21 and arranged according to the retention time (Rt), Supplementary Table S1]. The positive mode ESI-MS analysis of each collected fraction obtained from the P1 crude extract revealed a profile with major molecular ions at *m/z* 430.1609, 324.2073, 515.3331, 557.3804, 541.3485, 543.3644, 571.3931, and 559.3953 (compounds 1, 2, 4, 7, 9, 13, 18, and 14, respectively). The ESI-MS analysis of the NP1 crude extract revealed a similar profile with the major molecular ions of *m/z* 430.1609, 515.3331, 557.3804, 541.3485, 543.3644, 559.3953, and 585.4117 (compounds 1, 4, 7, 9, 13, 14, and 16, respectively). The corresponding sodium and potassium adducts of the compounds from the P1 and NP1 fractions were also detected (Supplementary Table S1). It should be noted that several compounds co-eluted with the major compounds (higher molecular ion signal) (Supplementary Table S1). Due to the co-elution of presumptive known (*n* = 12) and unknown (*n* = 9) compounds detected in the P1 and/or NP1 crude extracts, UPLC-MS^n^ coupled with molecular networking analysis was used to cluster structurally similar compounds and elucidate the putative structures of the unknown compounds and facilitate the confirmation of the known structures.

**Molecular Networking Analysis**

The fractions obtained after RP-HPLC of the secondary metabolites extracted from the P1 and NP1 strains were subjected to UPLC-MS^n^ analysis and the raw MS^n^ data was used to generate a molecular network. Analysis of the molecular network revealed four clusters (Figures 2, 3, 5), including families of serratiochelin (comprised of five nodes) and prodigiosin (comprised of two nodes) that were identified using the GNPS and MS-DIAL libraries, respectively, and two unidentified F1 (comprised of 15 nodes) and F2 (comprised of 22
nodes) families that did not correspond to compounds within the GNPS and MS-DIAL libraries. In general, the majority of the metabolites (nodes) clustered in the F1, F2 and serratiochelin families and were detected in both P1 and NP1 strains; however, only the P1 strain produced compounds in the prodigiosin cluster. The entire network was formed by 55 nodes, including 11 individual nodes.

**UPLC-MS² Analysis**

Following molecular networking, the structural elucidation of the four distinct metabolite clusters produced by the *S. marcescens* P1 and NP1 strains was conducted by analyzing the fragmentation profiles of the UPLC-MS² data. The identified serratiochelin cluster ([Figure 2](#fig2)) revealed a node with m/z 430.1609 [C₂₁H₂₃N₃O₇ + H]⁺ (compound 1) corresponding to the peak observed at 3.76 min in the UPLC-MS² data for the P1 and NP1 strains ([Table 1](#tab1); [Supplementary Figure S1](#supfig1)). The fragmentation pattern of 1, which was observed using UPLC-MS², corresponded to the serratiochelin A (also referred to as serranticin) structure ([Kuo et al., 2011; Seyedsayamdost et al., 2012](#ref1)). The prodigiosin cluster ([Figure 2](#fig2)) revealed a node with m/z 324.2073 [C₂₀H₂₅N₃O⁺ H]⁺ (compound 2) corresponding to the peak observed at 8.48 min in the UPLC-MS² data for the P1 strain ([Table 1](#tab1); [Supplementary Figure S2](#supfig2)). The fragmentation pattern of 2, which was observed using UPLC-MS², corresponded to the prodigiosin structure ([Lee et al., 2011; Su et al., 2016](#ref2)). The fragmentation profiles and structural elucidation of the compounds detected in clusters F1 and F2 will be discussed according to each metabolite family (cluster). A summary of the UPLC-MS² data for the metabolites within these clusters produced by the P1 and NP1 strains is presented in [Tables 1, 2](#tab2) and [Supplementary Figures S22, S23](#supfig22, #supfig23), and the structures of the known metabolites and putative structures of the unknown compounds produced by the P1 and NP1 strains are depicted in [Figures 3, 5](#fig3, #fig5) and [Supplementary Figures S3–S21](#supfig3–#supfig21).
Structural Elucidation of Serratamolide Homologues

The molecular network of the F1 cluster and corresponding structures for the known and novel compounds in the F1 cluster are presented in Figure 3. The F1 cluster (Figure 3) revealed nodes with m/z 515.3331 [C_{26}H_{46}N_{2}O_{8} + H]^{+} (compound 4), 541.3485 [C_{28}H_{48}N_{2}O_{8} + H]^{+} (compound 9), 543.3644 [C_{28}H_{50}N_{2}O_{8} + H]^{+} (compound 13), 561.3749 [C_{28}H_{52}N_{2}O_{9} + H]^{+} (compound 6), 559.3600 [C_{28}H_{50}N_{2}O_{9} + H]^{+} (compound 3), and 571.3931 [C_{30}H_{54}N_{2}O_{8} + H]^{+} (compound 18), which corresponded to peaks observed in the UPLC-MSE data for the P1 and NP1 strains (Table 1; Supplementary Figures S3–8, S22). In addition, the F1 cluster (Figure 3) revealed a node with m/z 587.3908 [C_{30}H_{54}N_{2}O_{8} + H]^{+} (compound 11) corresponding to the peak observed at 13.76 min in the UPLC-MSE data for the NP1 strain (Table 1; Supplementary Figures S9, S22). The fragmentation profiles of these seven compounds (3, 4, 6, 9, 11, 13, and 18) correspond to cyclic or open-ring serratamolide homologues that have previously been reported (Dwivedi et al., 2008; Eckelmann et al., 2018).

Compound 4 was thus comprised of two serine residues (cyclic peptide moiety) coupled to two saturated hydroxydecanoic acyl groups (C_{10}) (Figure 3) confirming the identity of the compound as serratamolide A (Dwivedi et al., 2008). In addition, 18 was comprised of two serine residues (cyclic peptide moiety) linked to two saturated hydroxydecanoic acyl groups (C_{12}), which corresponds to a serratamolide homologue reported by Eckelmann et al. (2018) (Figure 3). In contrast, 9 and 13 were comprised of two serine residues (cyclic peptide moiety) coupled to a 3-hydroxydecanoic acyl (C_{10}) and a hydroxydodecanoic acyl (C_{12}) (Figure 3). Compound 13 thus corresponded to the previously described serratamolide C (Dwivedi et al., 2008). However, compound 9 had a double bond in the hydroxydodecanoic acyl (C_{12,1}) residue (Figure 3) confirming the structure as serratamolide B (Dwivedi et al., 2008). The double bond position was putatively placed at the C_{5} position (from the C=O group) of the fatty acyl chain, as this corresponded to the
position previously reported in literature (Eckelmann et al., 2018). Compounds 3 and 6 had identical fatty acyl chain lengths as serratamolide B and C, respectively, however, the addition of a hydroxyl group, based on the experimental molecular formula (elemental composition) predicted using the MassLynx 4.1 software package, in the compound suggested that both lipopeptides were open-ring structures and corresponded to the fragmentation pattern of the previously described open-ring serratamolides B and C (Eckelmann et al., 2018) (Figure 3). Similarly, 11 was comprised of one serine residue coupled to one saturated hydroxydodecanoic acyl (C_{12}) and one serine residue coupled to one unsaturated hydroxydodecanoic acyl (C_{12:1}). However, the addition of a hydroxyl group [based on the experimental molecular formula (elemental composition)] in the compound suggested that the lipopeptide was an open-ring structure and corresponded to the fragmentation pattern previously described by Eckelmann et al. (2018) (Figure 3). Overall, it is proposed that this compound group is comprised of four residues, including a peptide moiety of two serine residues (cyclic or open-ring) attached to two fatty acyl residues (varying in length and presence or absence of double bonds). From the described serratamolides it was assumed that the compounds described above contain L-serine residues and D-3-hydroxy-fatty acyl residues.

The compound with m/z 585.374 [C_{30}H_{52}N_{2}O_{9} + H]^{+} (compound 8) did not cluster in the molecular network; however, it was detected at 12.46 min in the UPLC-MSE data for the NP1 strain (Table 1; Figure 4A; Supplementary Figures S10, S22). The fragment with m/z 567.368 corresponded to a product of dehydration (removal of H, OH). A major fragment at m/z 284.185 was then observed, representing a dehydrated serine residue linked to an unsaturated hydroxydodecanoic acid (C_{12:1}). In addition, the dehydration product of the m/z 284.185 fragment was detected at m/z 266.176. Similar to the previously described open-ring structures, the addition of a hydroxyl group [based on the experimental molecular formula (elemental composition)] in the compound suggested that the lipopeptide was an open-ring structure; however, this compound has not previously been reported in literature or databases (Norine and PubChem). The novel 8 homologue was thus proposed to comprised of two L-serine residues coupled to two unsaturated

![FIGURE 4](image-url)
D-3-hydroxydodecanoic acyl groups (C\(_{12:1}\)), correlating to described serratamolides (Figure 4A).

The F1 nodes with \(m/z\) 537.3147, 563.3290, 565.3455, 581.3405 and 609.3735 were found in the UPLC-MS\(^E\) data and corresponds to the sodiated adducts of the compounds 4, 9, 13, 14, 3, and 11, respectively, (Supplementary Table S1). In contrast, no structures were established for the F1 nodes with \(m/z\) 531.365 and 575.391.

### Structural Elucidation of Glucosamine Derivatives

The molecular network of the F2 cluster and corresponding structures for the known and novel compounds in the F2 cluster are presented in Figure 5. The F2 cluster (Figure 5) revealed nodes with \(m/z\) 585.4117 \([\text{C}_{31}\text{H}_{56}\text{N}_2\text{O}_8 + \text{H}]^+\) (compound 16) and 559.3953 \([\text{C}_{29}\text{H}_{54}\text{N}_2\text{O}_8 + \text{H}]^+\) (compound 14) corresponding to the peaks observed in the UPLC-MS\(^E\) data for the P1 and NP1 strains, while \(m/z\) 573.4136 \([\text{C}_{30}\text{H}_{56}\text{N}_2\text{O}_8 + \text{H}]^+\) (compound 19) corresponded to a peak observed in the UPLC-MS\(^E\) data for the P1 strain (Table 2: Supplementary Figures S11–S13, S23). The fragmentation profiles of these three compounds (14, 16, and 19) correspond to glucosamine derivative homologues that have previously been reported (Dwivedi et al., 2008). For all three compounds, the two major fragments were linked via an anomeric C-O bond between the hexose and valine amino acid residues. Therefore, 16 was proposed to be comprised of a hexose residue linked to butyric acyl group, which in turn was coupled to a valine residue linked to an unsaturated C\(_{16:1}\) fatty acid chain correlating to a previously described compound, glucosamine derivative A (Dwivedi et al., 2008) (Figure 5). The hexose was assumed to be glucose and the double bond within the fatty acyl chain of glucosamine derivative A was assumed to be a cis double bond corresponding to the structure previously described in literature (Dwivedi et al., 2008). Compound 19 was also proposed to be comprised of a glucose residue linked to a butyric acyl group, coupled to a valine residue linked to a saturated C\(_{15}\) fatty acid chain, thus correlating with the structure of glucosamine derivative B as described by Dwivedi et al., 2008.
et al. (2008) (Figure 5). Finally, 14 had a structure similar to the other two glucosamine derivatives. Compound 14 was also proposed to be comprised of a glucose residue linked to butyric acid, a valine residue linked to a saturated C14 fatty acid chain and thus the structure correlated to the glucosamine derivative C (Dwivedi et al., 2008) (Figure 5). Overall, it is proposed that this compound group is comprised of four residues, namely glucose, valine, butyric acid and a saturated or unsaturated β-fatty acid of varying length (C14 to C16).

Other F2 nodes (Figure 5) at m/z 557.3804 [C25H30N2O8 + H]+ (compound 7), 583.3947 [C31H32N2O8 + H]+ (compound 15) and 589.4056 [C30H56N2O9 + H]+ (compound 12) had similar fragmentation profiles to 14, 16, and 19 and corresponded to peaks observed in the UPLC-MSF data for P1 and NP1 strains (Table 2; Supplementary Figures S14–S16). In addition, F2 nodes at m/z 575.3902 [C29H34N2O9 + H]+ (compound 5), 587.4268 [C31H34N2O9 + H]+ (compound 21), and 599.4265 [C32H36N2O8 + H]+ (compound 17) also had similar profiles to glucosamine derivative A and C and corresponded to peaks observed in the UPLC-MSF data for the P1 strain, while another node at m/z 545.3787 [C28H32N2O9 + H]+ (compound 10) corresponded to peaks observed in the UPLC-MSF data for the NP1 strain (Table 2; Supplementary Figures S17–S20, S23). The cleavage of all seven compounds at the anomeric C-O bond resulted in two major fragments for each compound. Similar to the previously described known compounds 14, 16, and 19, each compound had a major fragment of m/z 232.116 corresponding to the glucose/hexose residue linked to butyric acid, and dehydration products of the m/z 232.116 fragment were observed at m/z 214.107 and 196.097. Compounds 10 and 21 had a second major fragment of m/z 314.272 and 356.316, which corresponded to a valine residue linked to a saturated C13 and C16 fatty acyl chain, respectively. In addition, 7, 15, and 17 had a second major fragment of m/z 326.267, 352.282, and 368.316 which corresponded to a valine residue linked to an unsaturated C14, C18:2 and C17:1 fatty acid chain, respectively. Finally, 5 and 12 had a major fragment of m/z 344.280 and 358.296, which corresponded to a valine residue linked to a saturated C14 and C15 fatty acyl chain with an additional hydroxyl group in the fatty acyl residue. The presence of the hydroxyl group within the fatty acyl moiety was further confirmed, as the loss of a hydroxyl group was observed within the fragmentation profile for both m/z 344.280 and 358.296 fragments, including m/z 326.269 and 340.286, respectively. In addition, the experimental molecular formula (elemental composition) predicted by the MassLynx 4.1 software package provided further indication of the addition of the hydroxyl group for both 5 and 12 (Table 2).
Typically, MS analysis is used in conjunction with other analytical methods, such as NMR, to determine the position of double bonds and hydroxyl groups within a structure (Eckelmann et al., 2018). However, the majority of the glucosamine derivative compounds were obtained in low quantities and could thus not be subjected to NMR analysis. A cis configuration was thus proposed for the double bonds in the fatty acid chain of the various unknown compounds, as the geometric isomerism of a double bond is typically a cis configuration in natural fatty acids (Kobelnik et al., 2018). Therefore, we propose that 7, 10, 12, 15, 17, and 21) and fragmentation patterns did not correspond to previously identified glucosamine derivatives (N-butylglucosamine ester derivatives) and are hypothesized to be novel glucosamine derivatives homologues. The F2 nodes with m/z 607.3924, 581.3781, 579.3623, and 567.3618 were found in the UPLC-MS² data and corresponded to the sodiated adducts of compounds 16, 14, 7, and 10, respectively, (Supplementary Table S1). No structures were elucidated for the F2 nodes with m/z 513.316, 543.363, 555.367, 569.381, and 601.407.

In contrast, to the above mentioned glucosamine derivatives, an individual node (Figure 2) with m/z 627.4192 [C₃₃H₅₈N₂O₉ + H]⁺ (compound 20) exhibited a unique fragmentation profile and corresponded to a peak at 16.28 min observed in the UPLC-MS² data for the P1 strain (Table 2; Figure 4B; Supplementary Figures S21, S23). The fragmentation profile was unique as 20 did not have a major fragment of m/z 232.116 (corresponding to a hexose residue linked to butyric acyl group) that was common in all of the previously described glucosamine derivatives. Instead, a major fragment at m/z 274.129 was observed, which suggested a modification in the fragment corresponding to the hexose and butyric acyl moiety (Figure 4B). The major fragment at m/z 274.129 potentially corresponded to a hexose residue linked to an oxo-hexanoic acyl (rather than a butyric acyl), and a dehydration product of the m/z 274.129 fragment was observed at m/z 256.117. Furthermore, a second major

![Figure 5](image-url)
fragment of m/z 354.302 was observed, which corresponded to an unsaturated C16:1 fatty acid chain. In comparison to the glucosamine derivative A (16) which also had an unsaturated C16:1 fatty acid chain, this glucosamine derivative had an additional C2H3O based on the experimental molecular formula (elemental composition) predicted using the MassLynx 4.1 software package. It is thus feasible that oxo-hexanoic acid was incorporated into the glucosamine derivative instead of butyric acid. In addition, 20 did not cluster with the glucosamine derivatives in the molecular network; rather, the compound formed an individual cluster and this further suggests the reduced relatedness of this compound to the glucosamine derivatives. Although NMR is required to confirm the structure, the putative novel congeners of 20 was proposed to consist of glucose, oxo-hexanoic acid, valine and an unsaturated C16:1 fatty acid chain, which is distinctive from that of the previously described glucosamine derivatives (Figure 4B).

Antimicrobial Activity

The bioactivity of the secondary metabolites produced by the P1 and NP1 strains was then investigated using disc diffusion, MIC and MBC assays. All the fractions collected during RP-HPLC analysis were subjected to antimicrobial testing using a disc diffusion assay against E. faecalis S1. It was found that E. faecalis S1 was susceptible (diameter range of 9.3 ± 2.1 to 19.3 ± 1.5 mm) to the fractions corresponding to the major compounds (higher molecular ion signal) 4, 9, 13, 7, 14, and 16 (for both P1 and NP1), and the fraction of compound 2 collected from P1 crude extract (results not shown). Thereafter, the MIC and MBC assays were performed using five fractions (selected based on activity and quantity available), including fractions of the major compounds (higher molecular ion signal) 2, 4, 9, 13, and 16. A summary of the MIC and MBC assay results are outlined in Table 3. It was observed that all three serratamolides (4, 9, and 13) (identified as outlined in Structural Elucidation of Serratamolide Homologues) exhibited identical activity against E. faecalis S1 with a MIC of 3 mg/ml and MBC of >3 mg/ml. In comparison, glucosamine derivative A (compound 16; identified as outlined in Structural Elucidation of Glucosamine Derivatives) was found to have a lower MIC and MBC of 0.75 mg/ml and 3 mg/ml, respectively. Overall, however, the lowest MIC and MBC recorded against E. faecalis S1 was observed for prodigiosin (compound 2; identified as outlined in UPLC-MS E Analysis), at <0.75 and 1.5 mg/ml, respectively, (Table 3).

DISCUSSION

Untargeted Metabolomics: UPLC-MS E and Molecular Networking Analysis

This work reports on the identification of four molecular families of secondary metabolites associated with Serratia species, including serratiochelin A (1), prodigiosin (2), serratomolide homologues (3, 4, 6, 8, 9, 11, 13, and 18) and glucosamine derivative congeners (5, 7, 10, 12, 14–17, and 19–21) (refer to Figures 2–5). The structures of prodigiosin and serratiochelin A have previously been elucidated, where prodigiosin was described as a tripyrrole red pigment with an alkylation substituent (Figure 1A; Lee et al., 2011) and serratiochelin A was described as a low-molecular weight siderophore with high affinity for iron (Figure 1B; Khilyas et al., 2019). The structure of several of the serratomolides and glucosamine derivative metabolites detected in the current study has previously been elucidated, with serratomolides generally described as lipopeptides of two L-serine residues (cyclic or open-ring) linked to two fatty acid chains (Figure 1C) and glucosamine derivatives described as glucose amine consisting of glucose, valine, a fatty acid chain and butyric acid (Figure 1D; Dwivedi et al., 2008; Eckelmann et al., 2018). Moreover, based on their characteristic fragmentation patterns, as observed in the UPLC-MS E data, eight serratomolides (including the putative structure of a novel open-ring serratomolide homologue) and 11 glucosamine derivatives (including the putative structures of eight novel glucosamine derivative congeners) were identified in this study and their structures were elucidated.

It is well-known that certain bacteria naturally produce various lipopeptide congeners which vary in fatty acid chain length, fatty acyl saturation or in the amino acyl residues of the peptide moiety (Das et al., 2008). This was similarly observed in the current study as the identified serratomolides, comprised of two L-serine residues linked to two fatty acyl residues, varied in the length (C10 to C12) and the presence or absence of double bonds (saturated or mono-unsaturated) in the fatty acid chains and cyclization (open-ring or cyclic) of the peptide moiety. The glucosamine derivative congeners, comprised of glucose, valine, butyric acyl (or oxo-hexanoic acyl for 20) and a fatty acyl residue, similarly varied in the length (C13 to C17), as well as the presence or absence of double bonds (saturated or unsaturated) or a hydroxyl group in the β-fatty acyl residue. During UPLC-MS E analysis, the elution time (Rt) of the serratomolide and glucosamine derivative homologues was, as expected,

### Table 3 | Minimum Inhibition and Bactericidal Concentrations of Selected Fractions Against the Clinical E. faecalis S1 Stain.

| Compound No | Proposed Identification (Fatty Acid Chain Length) | Purity | MIC (mg/ml) | MBC (mg/ml) |
|-------------|--------------------------------------------------|--------|-------------|-------------|
| 4           | Serratamolide A (C10 + C10)                       | 72%    | 3           | >3          |
| 9           | Serratamolide B (C10 + C12:1)                     | 61%    | 3           | >5          |
| 13          | Serratamolide C (C10 + C12)                       | 75%    | 3           | >3          |
| 16          | Glucosamine derivative A (C16:1)                  | 80%    | 0.75        | 3           |
| 2           | Prodigiosin                                       | 80%    | <0.75       | 1.5         |
dependent on the length of the β-hydroxy fatty acid chains of the compound. For instance, the serratamolides with shorter saturated fatty acid moieties, such as serratamolide A (C_{10} + C_{10}) (4), eluted from the column before the serratamolides with longer saturated fatty acid moieties, such as serratamolide 571 (C_{12} + C_{12}) (18) (refer to Table 1; Supplementary Figure S22). It was also observed that the Rt of the serratamolide and glucosamine derivative homologues was dependent on the saturation or unsaturation of the fatty acid moieties, for instance, as the unsaturated glucosamine derivative A (16) (C_{16}l) eluted before the corresponding saturated novel glucosamine derivative K (21) homologue (C_{16}S) with the same residues and fatty acid chain length (refer to Table 2; Supplementary Figure S23). Previous research has indicated that the presence and position of a double bond within the fatty acyl chain influences the Rt of a compound (Takashima et al., 2018) and the retention time of a mono-unsaturated fatty acid is reduced to that of the equivalent saturated fatty acid with two carbon units. A cis double bond causes a kink in the chain and reduces the actual molecular length by nearly two carbon units, thereby resulting in the elution of a C_{18} fatty acid just after a C_{16} fatty acid (Plattner et al., 1977; Takashima et al., 2018). It has also been reported that an open-chain lipopeptide is retained in the column for a shorter time period than the corresponding cyclic lipopeptide due to the increased polarity of the compound (due to the addition of the terminal COOH group) (Qiu et al., 2019), which clarifies for example, why the cyclic serratamolide C (C_{10} + C_{12}) (13) eluted from the column after the open-chain serratamolide 561 (C_{10} + C_{12}) (6) with the same fatty acid chain length. Furthermore, it was observed that the glucosamine derivatives with an additional hydroxyl group within the fatty acyl moieties were retained in the column for a shorter time period than the corresponding glucosamine derivative without a hydroxyl group, due to the increased polarity of the compound. For instance, the glucosamine derivative C (C_{29}H_{52}N_{2}O_{6}; 14) with a C_{14} fatty acyl chain eluted from the column after the corresponding novel glucosamine derivative D homologue (C_{29}H_{53}N_{2}O_{6}; 5) with a C_{14} fatty acyl chain that has an additional hydroxyl group. Overall, it was observed that the serratamolide and glucosamine derivative homologues separated as expected according to hydrophobicity (fatty acyl chain length) and chain saturation.

**Secondary Metabolic Profile of the S. marcescens P1 and NP1 Strains**

Clements et al. (2019b) previously identified the secondary metabolites produced by the P1 and NP1 strains using UPLC-MS analysis, which proposed that the P1 strain produced prodigiosin and serratamolides (A, B, C, and E), while the NP1 strain produced glucosamine derivative A and serratamolides (A, B, C, and E). The use of RP-HPLC and UPLC-MS analysis in the current study revealed that the P1 strain produced four metabolic classes, including prodigiosin, serratiochelin A, serratamolide homologues (n = 6) and glucosamine derivative congeners (n = 10), while NP1 produced three metabolic classes, including serratiochelin A, serratamolide homologues (n = 8) and glucosamine derivative congeners (n = 6). It is hypothesized that the combination of RP-HPLC and UPLC-MS analysis was an effective approach to identify the minor metabolic constituents and elucidate the microbial metabolic profile of the environmental *Serratia* strains.

Genetically, secondary metabolites are synthesized by a number of gene clusters that can be identified using a bioinformatics approach (Romano et al., 2018). Certain gene clusters involved in secondary metabolism [such as non-ribosomal peptide synthetase (NRPS) gene clusters] may not be expressed under laboratory conditions and may be considered as “silent” gene clusters (Bode et al., 2002). Minor alterations in the cultivation conditions of a particular strain may, however, trigger the expression of the “silent” gene clusters and may allow for the discovery of novel small molecules (Bode et al., 2002). This phenomenon is known as the “one strain many compounds” (OSMAC) approach, which indicates that a single strain can produce various metabolites when grown under different media composition and cultivation conditions, leading to the activation of the “silent” metabolic pathways within the genome of the microbial strain (Bode et al., 2002; Romano et al., 2018). When comparing the culture conditions of Clements et al. (2019b) to the culture conditions employed in the current study, an increased production scale (from 100 ml to 1,500 ml) in baffled flasks, as well as the change in media components (from bactopeptone to peptone powder) was applied. Thus, while the combination of RP-HPLC and UPLC-MS analysis may have contributed to the detection of the minor metabolites; it is hypothesized that the altered culture conditions and media composition employed, may have activated “silent” NRPS clusters (such as the glucosamine derivatives in the P1 strain and the serratiochelin in both strains), thereby increasing the metabolic constituents produced by both the pigmented and non-pigmented *S. marcescens* strains.

Although genome mining of the P1 and NP1 strains was not within the scope of this study, Khilyas et al. (2019) sequenced the genome of a pigmented *S. marcescens* SM6 strain and used the antiSMASH 5.0 prediction tool to identify biosynthetic gene clusters involved in secondary metabolism within the strain. The authors identified five biosynthetic gene clusters involved in secondary metabolite production, including gene clusters responsible for the production of two siderophores (serratiochelin and chrysobactin), a thiopeptide, a serratamolide and a glucosamine derivative (Khilyas et al., 2019). It is possible that the P1 and NP1 strains possess similar NRPS gene clusters as both strains were able to produce serratiochelin, serratamolides and glucosamine derivatives. In addition, the bifurcated biosynthetic pathway responsible for the production of prodigiosin was additionally activated in the P1 strain (Lee et al., 2011). Following an extensive literature search, and to the best of the author’s knowledge, this is the first report of the co-production of all four secondary metabolic classes (including prodigiosin, serratiochelin A, serratamolides and glucosamine derivatives) by a *Serratia* strain.
Antimicrobial Activity

Results from this study indicated that prodigiosin (2), serratamolide A (4), B (9), and C (13) homologues and glucosamine derivative A (16) and C (14) homologues, as well as the novel glucosamine derivative E (7) homologue exhibited activity against a clinical Gram-positive bacterial strain. The MIC and MBC assays further revealed that serratamolide A (4), B (9), and C (13) exhibited identical activity against E. faecalis S1, suggesting that the presence of a double bond or the increased length of one of the two fatty acid chains from C_{10} to C_{12} exhibited no additional antimicrobial potency against the Gram-positive strain. In comparison, glucosamine derivative A (16) was found to exhibit a lower MIC and MBC in comparison to the serratamolides. This indicates that E. faecalis S1 was more susceptible to the glucosamine derivative A (16) than the serratamolides. Overall, however, E. faecalis S1 was found to be the most susceptible to prodigiosin (2), as a lower MIC and MBC was obtained against the clinical strain in comparison to glucosamine derivative A (16) and the serratamolides (4, 9, and 13).

The potent activity of prodigiosin corresponds to previous research that similarly recorded activity of this pigmented compound against Gram-positive bacteria such as Staphylococcus aureus (S. aureus) and Bacillus subtilis (B. subtilis) (Darshan and Manonmani, 2015). It has also been suggested that prodigiosin is a hydrophobic stressor that disrupts the cell membrane and results in the leakage of intracellular products, such as K⁺ ions, amino acids, sugars, and proteins (Suryawanshi et al., 2016). Furthermore, prodigiosin may interfere with cellular respiration and protein and ribosomal ribonucleic acid synthesis (Danevčić et al., 2016a; Danevcic et al., 2016b). Serratamolide A has similarly been reported to display activity toward Gram-positive bacteria, such as S. aureus strains (Kadouri and Shanks, 2013). It has additionally been reported that serratamolide A (serrawettin W1) increased the rate of movement of K⁺ and H⁺ across the membrane of S. aureus at a concentration of 10 μg/ml, while no leakage of “260 nm-absorbing products” (such as nucleic acids), membrane permeability or cell growth inhibition was observed at this concentration (Deol et al., 1973). Currently, one study has reported on the antimicrobial activity of glucosamine derivative homologues (A to C) toward Mycobacterium diernhoferi; however, little is known about the mechanism of action of these metabolites (Dwivedi et al., 2008). Moreover, Hage-Hülsmann et al. (2018) described the enhanced antimicrobial activity of serrawettin W1 (serratamolide A) and prodigiosin in combination, in comparison to the individual bioactive metabolites, against Corynebacterium glutamicum. The authors hypothesized that the co-production of serrawettin W1 and prodigiosin may provide a competitive advantage to the producing strain due to the increased bioactive potency, while also aiding in the colonization and biofilm development of the producing strain by inhibiting or delaying the growth of competing microorganisms. Thus, while extensive research is required to investigate the ecological role of co-producing prodigiosin, serratamolides and glucosamine derivatives; the combined production of these bioactive metabolites by S. marcescens may provide a competitive advantage to the producing strain (Darshan and Manonmani, 2015; Hage-Hülsmann et al., 2018). Moreover, as indicated in Clements et al. (2019b), the metabolites in the P1 strain may exhibit a higher antimicrobial effect in comparison to the metabolites in the NP1 strain, as the secondary metabolites in the P1 crude extract were effective against 81% of the test microbial organisms, in comparison to the 67% antimicrobial efficacy recorded for the NP1 strain.

CONCLUSION

The use of RP-HPLC for chromatographic fractionation, high resolution ESI-MS, UPLC-MS² and molecular networking analysis allowed for the identification and elucidation of 21 secondary metabolites (combined total for P1 and NP1 strains), including serratamolide homologues, glucosamine derivative homologues, prodigiosin (P1) and serratiocelin A. Moreover, the putative structure of an open-ring serratamolide homologue and putative structures of eight novel glucosamine derivative congeners were elucidated. The structural relatedness of the novel open-ring serratamolide and glucosamine derivative congeners to the known serratamolides and glucosamine derivatives, respectively, was revealed using molecular networking and fragmentation patterns of the UPLC-MS² data. This approach thus proved vital for metabolite identification and characterization. The identified metabolic families exhibited structural diversity and well-ordered chromatographic behaviors due to structural modifications of fatty acid chain lengths, open-ring or cyclic structures and the degree of saturation (saturated or mono-unsaturated) of the fatty acid chain. Moreover, seven fractions collected during RP-HPLC, with major molecular ions corresponding to prodigiosin (2), serratamolide A (4), B (9), and C (13) homologues and glucosamine derivative A (16), C (14), and E (7) homologues, displayed antimicrobial activity against the clinical E. faecalis S1 strain using the disc diffusion assay. However, the MIC and MBC assays revealed that the three serratamolides [A (4), B (9), and C (13)] exhibited the lowest antimicrobial potency (same MIC and MBC values for all three) of the metabolites tested, and highlighted that presence of a double bond or the change in length of a fatty acid chain (C10 to C12) of the serratamolides provided no additional antimicrobial potency against the Gram-positive strain. Prodigiosin (2) however, exhibited the greatest antimicrobial potency, followed by glucosamine derivative A (16). This study thus highlights the importance of Serratia species as a source of novel antimicrobial compounds for potential therapeutic application. It is recommended that future research investigate the mode of action and synergism of the bioactive secondary metabolites produced by these two S. marcescens strains. In addition, the use of whole genome sequencing of the P1 and NP1 strains and subsequent genome mining of NRPS gene clusters is a promising future endeavour to identify additional secondary metabolites not actively being produced under typical cultivation conditions.
DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

Conceived and designed the experiments: TC and WK. Performed the experiments: TC. Analyzed the data: TC and MR. Contributed reagents/materials/analysis tools: WK and SK. Compiled the manuscript: TC and WK. Edited the manuscript: MR, SK, and TN.

FUNDING

This work was financially supported by the Water Research Commission (grant number: K5/2728/[3]) and the National Research Foundation of South Africa (grant number: 113849).

REFERENCES

Bode, H. B., Bethe, B., Hofs, R., and Zeeck, A. (2002). Big effects from small changes: possible ways to explore nature’s chemical diversity. ChemBioChem 3 (7), 619–627. doi:10.1002/1439-7633(20020703)3:7<619:AID-CBIC619>3.0.CO;2-9
Clements, T., Ndlovu, T., Khan, S., and Khan, W. (2019a). Biosurfactants produced by endophytic Serratia species: classification, biosynthesis, production and application. Appl. Microbiol. Biotechnol. 103, 589–602. doi:10.1007/s00253-018-9520-5
Clements, T., Ndlovu, T., and Khan, W. (2019b). Broad-spectrum antimicrobial activity of secondary metabolites produced by Serratia marcescens strains. Microbiol. Res. 229, 126329. doi:10.1016/j.micres.2019.126329
Danević, T., Vežjak, M. B., Tabor, M., Zorec, M., and Stopar, D. (2016a). Prodigiosin induces autolysis in actively grown Bacillus subtilis cells. Front. Microbiol. 7, 27. doi:10.3389/fmicb.2016.00027
Danević, T., Vežjak, M. B., Zorec, M., and Stopar, D. (2016b). Prodigiosin—a multifaceted Escherichia coli antimicrobial agent. PLoS One 11 (9), 1–13. doi:10.1371/journal.pone.0162412
Darshany, N., and Manonmani, H. K. (2015). Prodigiosin and its potential applications. J. Food Sci. Technol. 52, 5393–5407. doi:10.1007/s13197-015-1740-4
Das, P., Mukherjee, S., and Sen, R. (2008). Antimicrobial potential of a lipopeptide biosurfactant derived from a marine Bacillus circulans. J. Appl. Microbiol. 104 (6), 1675–1684. doi:10.1111/j.1365-2672.2007.03701.x
Deol, B. S., Bermingham, M. A., Still, J. L., Haydon, D. A., and Gale, E. F. (1973). The action of serratamolide on ion movement in lipid bilayers and biomembranes. Biochim. Biophys. Acta Biomembr. 330 (2), 192–195. doi:10.1016/0005-2736(73)90224-1
Dwivedi, D., Jansen, R., Molinari, G., Nimtz, M., Johri, B. N., and Wray, V. (2008). Antimycobacterial serratamolides and diacyl peptoglucosamine derivatives from Serratia sp. J. Nat. Prod. 71 (4), 637–641. doi:10.1021/jp070126
Eckelmann, D., Spiteller, M., and Kusari, S. (2018). Spatial-temporal profiling of prodigines and serratamolides produced by endophytic Serratia marcescens harbored in Maytenus serrata. Sci. Rep. 8, 1–15. doi:10.1038/s41598-018-23538-5
European Committee on Antimicrobial Susceptibility Testing. (2018). Breakpoint tables for interpretation of MICs and zone diameters. Available at: http://www.eucast.org/fileadmin/src/media/PDFs/EUCAST_files/Breakpoint_tables/v_8.0_Breakpoint_Tables.pdf (Accessed July 10, 2020).

Opinions expressed and conclusions arrived at, are those of the authors and are not necessarily to be attributed to the National Research Foundation.

ACKNOWLEDGMENTS

The authors wish to thank ADP van Staden at Stellenbosch University for assistance with the RP-HPLC analysis to purify the crude extracts. The authors also thank M Stander from the LCMS unit at the Central Analytical Facility, Stellenbosch University, for assistance with the mass spectrometric protocols and sample analyses used in this study.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2021.633870/full#supplementary-material.

Fu, Y., Luo, J., Qin, J., and Yang, M. (2019). Screening techniques for the identification of bioactive compounds in natural products. J. Pharm. Biomed. Anal. 168, 189–200. doi:10.1016/j.jpba.2019.02.027
Grumont, F., and Grumont, P. A. D. (2015). “Serratia,” in Bergey’s manual of systematicsof archaea and bacteria. Editors P. DeVos, J. Chun, S. Dedeyn, B. Hedlund, P. Kämpfer, F. Raine, et al. ( Hoboken, NJ: John Wiley & Sons), 1–81.
Hage-Hülsmann, J., Grünerberger, A., Thies, S., Santiago-Schübel, B., Klein, A. S., Petruzska, J., et al. (2018). Natural biocide cocktails: combinatorial antibiotic effects of prodigiosins and biosurfactants. PLoS One 13 (7), e0200940. doi:10.1371/journal.pone.0200940
Kadouri, D. E., and Shanks, R. M. (2013). Identification of a methicillin-resistant Staphylococcus aureus inhibitory compound isolated from Serratia marcescens. Res. Microbiol. 164 (8), 821–826. doi:10.1016/j.resmic.2013.06.002
Khlíyas, I. V., Tursunov, K. A., Shishrikova, T. V., Kamaletdinova, L. K., Matrosova, L. E., Desai, P. T., et al. (2019). Genome sequence of pigmented siderophore-producing strain Serratia marcescens SM6. Microbiol. Resour. Announc. 8 (18), e00247–19. doi:10.1128/MRA.00247–19
Knollhoff, A. M., Zheng, J., McFarland, M. A., Luo, Y., Callahan, J. H., Brown, E. W., et al. (2015). Identification and structural characterization of naturally-occurring broad-spectrum cyclic antibiotics isolated from Paenibacillus. J. Am. Soc. Mass Spectr. 26 (10), 1768–1779. doi:10.1007/s13136-015-1190-2
Kobelnik, M., Fontanari, G. G., Ribeiro, C. A., and Crespi, M. S. (2018). Evaluation of thermal behavior and chromatographic characterization of oil extracted from seed of Pittosporum undulatum. J. Therm. Anal. Calorim. 131 (1), 371–378. doi:10.1007/s10973-017-7653-9
Kuo, Y. H., Liang, T. W., Liu, K. C., Hsu, Y. W., Hsu, H. C., and Wang, S. L. (2011). Isolation and identification of a novel antioxidant with antitumour activity from Serratia ureilytica using squid pen as fermentation substrate. Mar. Biotechnol. 13 (3), 451–461. doi:10.1007/s10126-010-9316-9
Lee, J. S., Kim, Y. S., Park, S., Kim, J., Kang, S. J., Lee, M. H., et al. (2011). Exceptional production of both prodigiosin and cycloprodigiosin as major metabolic constituents by a novel marine bacterium, Zoohikella rubida S1-1. Appl. Environ. Microbiol. 77 (14), 4967–4973. doi:10.1128/AEM.01986-10
Magalhães, L., and Nitschke, M. (2013). Natural biocide cocktails: combinatorial antibiotic effects of prodigiosins and biosurfactants. PLoS One 13 (7), e0200940. doi:10.1371/journal.pone.0200940
Matysuama, T., Fujita, M., and Yano, I. (1985). Wetting agent produced by Matsuyama, T., Fujita, M., and Yano, I. (1985). Wetting agent produced by Matsuyama, T., Fujita, M., and Yano, I. (1985). Wetting agent produced by Matsuyama, T., Fujita, M., and Yano, I. (1985). Wetting agent produced by Matsuyama, T., Fujita, M., and Yano, I. (1985). Wetting agent produced
Matsuyama, T., Kaneda, K., Nakagawa, Y., Isa, K., Hara-Hotta, H., and Yano, I. (1992). A novel extracellular cyclic lipopeptide which promotes flagellum-dependent and -independent spreading growth of *Serratia marcescens*. *J. Bacteriol.* 174 (6), 1769–1776. doi:10.1128/jb.174.6.1769-1776.1992

Matsuyama, T., Murakami, T., Fujita, M., Fujita, S., and Yano, I. (1996). Extracellular vesicle formation and biosurfactant production by *Serratia marcescens*. *Microbiol. 132* (4), 865–875. doi:10.1099/00221287-132-4-865

Motley, J. L., Stamps, B. W., Mitchell, C. A., Thompson, A. T., Cross, J., You, J., et al. (2016). Opportunistic sampling of roadkill as an entry point to accessing natural products assembled by bacteria associated with non-anthopoid mammalian microbiomes. *J. Nat. Prod.* 80, 598–608. doi:10.1021/acs.jnatprod.6b00772

Ndlouvo, T., Rautenbach, M., Vosloo, J. A., Khan, S., and Khan, W. (2017). Characterisation and antimicrobial activity of biosurfactant extracts produced by *Bacillus amyloliquefaciens* and *Pseudomonas aeruginosa* isolated from a wastewater treatment plant. *AMB Express 7* (108), 1–19. doi:10.1186/s13568-017-0363-8

Philippus, A. C., Zatelli, G. A., Wanke, T., Barros, M. G. D. A., Kami, S. A., Lhullier, C., et al. (2018). Molecular networking prospection and characterization of terpenoids and C15-acetogenins in Brazilian seaweed extracts. *RSC Adv.* 8 (52), 29654–29661. doi:10.1039/C8RA02802H

Plattner, R. D., Spencer, G. F., and Kleiman, R. (1977). Triglyceride separation by reverse phase high performance liquid chromatography. *J. Am. Oil Chem. Soc.* 54 (11), 511–515. doi:10.1139/oil-77-11-0511

Qiu, S., Avula, B., Guan, S., Ravu, R. R., Wang, M., Zhao, J., et al. (2019). Identification of fusaricidins from the antifungal microbial strain *Paenibacillus* sp. MS2379 using ultra-high performance liquid chromatography coupled to quadrupole time-of-flight mass spectrometry. *J. Chromatogr. A.* 1586, 91–100. doi:10.1016/j.chroma.2018.12.007

Romano, S., Jackson, S. A., Patry, S., and Dobson, A. D. (2018). Extending the “one strain many compounds” (OSMAC) principle to marine microorganisms. *Mar. Drugs 16* (7), 244. doi:10.3390/md16070244

Seyedsayamdost, M. R., Cleto, S., Carr, G., Vlamakis, H., João Vieira, M., Kolter, R., et al. (2012). Mixing and matching sidrophore clusters: structure and biosynthesis of serratichelins from *Serratia* sp. V4. *J. Am. Chem. Soc.* 134 (33), 13550–13553. doi:10.1021/ja304941d

Soenens, A., and Imperial, J. (2019). Biocontrol capabilities of the genus *Serratia*. *Phytochem. Rev.* 2019, 1–11. doi:10.1007/s11101-019-09657-5

Stankovic, N., Senerovic, L., Ilic-Tomic, T., Vasiljevic, B., and Nikodinovic-Runic, J. (2014). Properties and applications of undecylprodigiosin and other bacterial prodigiosins. *Appl. Microbiol. Biotechnol.* 98, 3841–3858. doi:10.1007/s00253-014-5970-1

Su, C., Xiang, Z., Liu, Y., Zhao, X., Sun, Y., Li, Z., et al. (2016). Analysis of the genomic sequences and metabolites of *Serratia surfactantfaciens* sp. nov. YD25 that simultaneously produces prodigiosin and serrawettin W2. *BMC Genom.* 17, 865. doi:10.1186/s12864-016-3171-7

Suryawanshi, R. K., Patil, C. D., Koli, S. H., Hallsworth, J. E., and Patil, S. V. (2016). Antimicrobial activity of prodigiosin is attributed to plasma membrane damage. *Nat. Prod. Res.* 31 (5), 572–577. doi:10.1080/14786419.2016.1193580

Takashima, S., Toyoshi, K., and Shimozawa, N. (2018). Analyses of the fatty acid separation principle using liquid chromatography–mass spectrometry. *Med. Mass Spectrom.* 2, 1–13. doi:10.24508/mms.2018.06.002

Thies, S., Santiago-Schübel, B., Kovačić, F., Rosenau, F., Hausmann, R., and Jaeger, K. E. (2014). Heterologous production of the lipopeptide biosurfactant serrawettin W1 in *Escherichia coli*. *J. Biotechnol.* 181, 27–30. doi:10.1016/j.jbiotec.2014.03.037

Tsugawa, H., Caijka, T., Kind, T., Ma, Y., Higgins, B., Ikeda, K., et al. (2015). MS-DIAL: data-independent MS/MS deconvolution for comprehensive metabolome analysis. *Nat. Methods 12* (6), 523–526. doi:10.1038/nmeth.3393

Wang, M., Carver, J. J., Phelan, V. V., Sanchez, L. M., Garg, N., Peng, Y., et al. (2016). Sharing and community curation of mass spectrometry data with global natural products social molecular networking. *Nat. Biotechnol.* 34 (8), 828–837. doi:10.1038/nbt.3597

Wasserman, H. H., Keggi, J. J., and McKeon, J. E. (1961). Serratamolide, a metabolite product of *Serratia*. *J. Am. Chem. Soc.* 83 (19), 4107–4108. doi:10.1021/ja1480a046

Williamson, N. R., Fineran, P. C., Leeper, F. J., and Salmond, G. P. (2006). The biosynthesis and regulation of bacterial prodiginines. *Nat. Rev. Microbiol.* 4 (12), 887–899. doi:10.1038/nrmicro1331

Yasir, M., Dutta, D., and Willcox, M. D. (2019). Comparative mode of action of the antimicrobial peptide melimine and its derivative Mel4 against *Pseudomonas aeruginosa*. *Sci. Rep.* 9 (1), 1–12. doi:10.1038/s41598-019-42440-2

Yip, C. H., Yarkoni, O., Ajikoka, J., Wan, K. L., and Nathan, S. (2019). Recent advancements in high-level synthesis of the promising clinical drug, prodigiosin. *Appl. Microbiol. Biotechnol.* 103 (4), 1667–1680. doi:10.1007/s00253-018-09611-z

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.