CORRECTION

Correction: Screening and purification of natural products from actinomycetes that affect the cell shape of fission yeast (doi:10.1242/jcs.194571)

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There was an error published in J. Cell Sci. (2017) 130, 3173-3185 (doi:10.1242/jcs.194571).

In the above paper, we described the purification of a polyene from strain IS1 and identified it as a novel compound based on mass spectrometry data (published in Fig. 8B). We have been investigating this molecule in more detail and now believe, based on $^1$H NMR data, that it is identical to the known molecule fungichromin (also known as cogomycin, pentamycin and lagosin). The original identification of the molecule as novel was based on our interpretation of the ion at $m/z=693.379$ as a protonated ion [M+H]$^+$, whereas we now believe that this represents a sodiated ion [M+Na]$^+$.

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The authors apologise to the readers for any confusion that these errors might have caused.
Screening and purification of natural products from actinomycetes that affect the cell shape of fission yeast

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ABSTRACT
This study was designed to identify bioactive compounds that alter the cellular shape of the fission yeast Schizosaccharomyces pombe by affecting functions involved in the cell cycle or cell morphogenesis. We used a multidrug-sensitive fission yeast strain, SAK950 to screen a library of 657 actinomycete bacteria and identified 242 strains that induced eight different major shape phenotypes in S. pombe. These include the typical cell cycle-related phenotype of elongated cells, and the cell morphology-related phenotype of rounded cells. As a proof of principle, we purified four of these activities, one of which is a novel compound and three that are previously known compounds, leptomycin B, streptonigrin and cycloheximide. In this study, we have also shown novel effects for two of these compounds, leptomycin B and cycloheximide. The identification of these four compounds and the explanation of the S. pombe phenotypes in terms of their known, or predicted bioactivities, confirm the effectiveness of this approach.

KEY WORDS: Fission yeast, Actinomycetes, Natural product, Cell shape, Polyene, Leptomycin B, Cycloheximide

INTRODUCTION
The fission yeast Schizosaccharomyces pombe is a rod-shaped unicellular eukaryote that grows by apical extension and divides by medial fission and septation (Mitchison and Nurse, 1985). It has a typical eukaryotic cell cycle and this, together with the highly polarised growth pattern, makes it an excellent organism for studying the mechanisms involved in cell reproduction and the generation of cell form. Both the cell cycle and cell morphology are frequently altered in cancer cells, and mutations that alter these processes play a key role in the development of disease, for example in altering growth control (Beauchamp and Platania, 2013; Johnson and Halder, 2014), promoting delamination from an epithelial layer (Dekanty et al., 2015) and penetration of surrounding tissues during metastasis (Broustas and Lieberman, 2011). Indeed, morphological features are frequently used in cancer diagnosis. Around 142 genes have been identified in S. pombe as being conserved in humans and involved in cancer (Wood et al., 2002; http://www.pombase.org). It follows that compounds capable of perturbing the cell cycle or cell morphology in S. pombe may have similar effects in human cells and thus find utility in understanding the cell cycle, cell morphology and cancer, and in the development of anticancer treatments.

Random or site-specific mutagenesis, followed by screening for cells exhibiting aberrant shapes has allowed identification of many genes involved in the cell cycle and cell morphology in fission yeast (Nurse et al., 1976; Nurse and Thuriaux, 1980; Nasmyth and Nurse, 1981; Verde et al., 1995; Snaith and Sawin, 2003). These approaches led to the identification of the conserved Cdc2 (CDK1 in mammals) kinase, its regulatory subunit, the cyclin B Cdc13, and its regulators, the Wee1 tyrosine kinase and the Cdc25 tyrosine phosphatase (Nurse, 1990). Dephosphorylation of Cdc2 at Y15 allows a rise in Cdc2 kinase activity and entry into mitosis; thus, its timing is critical for regulating entry into mitosis (Simanis and Nurse, 1986; Russell and Nurse, 1986, 1987; Gould and Nurse, 1989). Other approaches have utilised near genome-wide deletion collections of essential and/or non-essential genes to identify new genes involved in these processes (Deshpande et al., 2009; Kelly and Nurse, 2011; Navarro and Nurse, 2012; Hayles et al., 2013; Graml et al., 2014).

As an extension of these approaches, we have designed a screen for natural compounds having an effect on the cellular shape of S. pombe to identify chemical entities affecting the functions involved in these processes. The development and use of such compounds as research tools to dissect the molecular mechanisms regulating the cell cycle and cell morphology will increase the range of available experimental methodologies. We have previously screened libraries of synthetic compounds produced by combinatorial chemistry (Takemoto et al., 2016; Kawashima et al., 2016), and decided to extend these studies by screening a library of bacterial isolates comprised of actinomycete strains. Actinomycetes are well known for synthesising natural products, for example, bleomycin (Umeno et al., 1966) and adriamycin (Arcamone et al., 1969), with anti-eukaryotic cell activities. The rationale behind this approach was that as the mechanisms that determine cell morphology and cell cycle in prokaryotic and eukaryotic organisms are different, and as soil-dwelling bacteria may be competing in the same ecological niche as fungi, these eukaryotic-specific processes could be targets for bacterially synthesised secondary metabolites (Ho and Nodwell, 2016). Thus, evolution would have naturally selected for prokaryotes able to produce antifungal agents, and thus may provide a richer source of ‘hits’ than would be obtained from a library of randomly synthesised molecules.

Phenotypic screens to identify novel bioactive natural products have been used previously; for example, fumagillin was isolated from fungal metabolites, and this compound induced endothelial cell rounding and inhibited angiogenesis (Inger et al., 1990).
Additionally, cell cycle-blocking agents from actinomycetes and fungi have been identified using *S. cerevisiae* as a screening tool (Tsuchiya et al., 2010). A number of screens have also been carried out in *S. pombe* using the viable haploid gene deletion mutant library to examine sensitivity of these mutants to known compounds (Deshpande et al., 2009; Gatti et al., 2011), and a study using wild-type *S. pombe* has looked at the effects of known natural products on cell shape by using FACS (Heisler et al., 2014). The objective of the work described here is to identify bioactive natural products through their effects on *S. pombe* cell shape by visual microscopic screening. We have assayed 657 bacterial strains for their effects on the multidrug-sensitive fission yeast strain SAK950 (Table S1), and report further analysis of the induced phenotype and purification of the active compound from four of these strains as a proof of principle for this experimental approach.

**RESULTS**

**Cell shape screening assay**

To examine the effects of bioactivities, we used a *S. pombe* strain (SAK950) with increased sensitivity to different compounds due to deletion of seven protein-encoding genes affecting drug efflux (Aoi et al., 2014) (Table S2).

A total of 657 actinomycete bacterial strains (http://demuris.co.uk) were tested for the production of bioactivities affecting the cellular shape of fission yeast (see Materials and Methods for details). We found 420 bacterial strains that led to the formation of a halo around the bacterial plug, indicating the presence of inhibitory bioactive natural products produced by the bacterial strain. For the remaining strains, 237 did not form a halo, and the *S. pombe* strains in the vicinity of the plug showed a wild-type phenotype.

Microscopic examination of *S. pombe* cells showed that of the 420 actinomycete strains that produced a halo, 178 killed fission yeast without showing an obvious shape phenotype (described as ‘dead’ in Table S1). The remaining 242 strains produced activities that induced a range of altered *S. pombe* cell shape phenotypes. We classified these phenotypes into eight broad groups (Fig. 1, Tables S3 and S4), related to the phenotypes observed in the deletion mutant collection screen (Hayles et al., 2013). However, the range of phenotypes exhibited did not completely overlap; for example, analysis of the gene deletion collection did not identify the swollen cell tip phenotype shown in Fig. 1C, suggesting that bioactive compounds may induce a different range of phenotypes compared with the gene deletions. This may be, for example, because they can interact with several different target molecules/binding sites within the cell, selectively inhibit particular activities of multifunctional target molecule, or can modify, rather than inactivate, a target molecule. We found that several strains induced mixed phenotypes, for example, both ‘small’ and ‘rounded’, or both ‘asymmetric septum’ and ‘misshapen’; however, for the purposes of classification, each of the bacterial strains was allocated to a single phenotypic group based on its predominant phenotypic effect.

From the 242 strains with activities, we selected 149 strains, which included representatives of all phenotypes classes (Tables S1 and S3), for further investigation due to the high penetrance or...
interest of the induced phenotypes. From this subset of strains, we selected activities produced from B21P2 (asymmetric septum), T342 (elongated), 5307(1B) (swollen cell tip) and IS1 (rounded) actinomycete strains for immediate further analysis to establish whether our approach was useful.

Characterization of an S. pombe asymmetry phenotype induced by B21P2

The asymmetric septation phenotype induced by B21P2 (Fig. 1J) was selected because this phenotype may be caused by a defect in spatial organisation within the cell. During interphase, microtubules are required for positioning of the nucleus in the middle of the cell, and the position of the septum at cytokinesis is normally dependent on the position of the nucleus (Chang and Nurse, 1996; Paoletti and Chang, 2000; Daga et al., 2006). It is therefore possible that this asymmetric septum phenotype induced by B21P2 is associated with other asymmetries and may be the result of a defective microtubule cytoskeleton and nuclear positioning.

We treated PN5352 cells (Table S2) growing in YE4S liquid medium with semi-purified extracts from B21P2 to examine the microtubule cytoskeleton and nuclear position. A range of phenotypes, including reduction or absence of interphase microtubules, and asymmetric positioning of the nuclei and septum (Fig. 2, see legend for details; Movies 1 and 2) was observed. Cells also had a range of spindle defects, with one of the predominant phenotypes observed being a nuclear microtubule bundle or extended spindle (Fig. 2; Movie 2). It is possible that this microtubule bundle/spindle may be related to the intranuclear bundle observed when Ned1 is overproduced (Tange et al., 2002). Formation of both the mitotic spindle and interphase microtubules is dependent on the import and export of tubulin and factors required for polymerization of microtubules (Sato and Toda, 2010). This may be inhibited in the presence of B21P2 and lead to microtubule cytoskeletal defects. In addition to microtubule defects, we also observed asymmetric division of the nuclear membrane in the presence of an abnormal mitotic spindle. It has previously been observed that the nuclear membrane can undergo fission in the absence of a mitotic spindle (Castagnetti et al., 2010). Our results suggest that in the presence of B21P2 there may be a defect in spindle attachment to the nuclear membrane, which allows membrane fission to occur independently of spindle elongation (Movie 2, lower right cell). B21P2 may also directly affect the nuclear membrane and lead to differently sized nuclei (Fig. 2; Movie 2, upper right cell). Some of these microtubule and nuclear membrane defects have previously been described for cells treated with leptomycin B (LMB), which inhibits nuclear export (Nishi et al., 1994). We tested the effect of LMB on PN5352 cells, and observed that the phenotype of SAK950 was almost identical to that seen with B21P2 (Fig. 2), suggesting the bioactive compound produced by this strain might be LMB.

Purification and identification of the bioactive compound from B21P2

To identify the natural product produced by B21P2 that is responsible for the asymmetric septum phenotype, we purified it and compared it with LMB. We found that medium D (INA5) was the most effective production medium for the bioactive compound after 7 days growth. Using these conditions, agar plates were harvested and the bioactive compound was extracted by reverse-phase flash chromatography, followed by an ethyl acetate extraction, normal-phase flash chromatography, size exclusion chromatography, preparative high-performance liquid chromatography (HPLC) and mass spectrometry, as described in the Materials and Methods.

The results of the purification indicated that there were two differing forms of the bioactive compound, which gave two discrete peaks in the HPLC elution at an λ=350 nm absorption spectrum trace. Eluate corresponding to both peaks was subjected to mass spectrometry, which revealed the presence of an ion at m/z=549.320 [M+Na]⁺ eluting at 14.6 min and an ion at m/z=563.331 [M+Na]⁺ eluting at 15.2 min (Fig. 3A,B shows the spectra from the two B21P2-derived active compounds). Quering of the ‘Dictionary of Natural Products’ (http://dnp.chemnetbase.com/faces/chemical/ChemicalSearch.xhtml) suggested that the most likely candidates were leptomycin A (LMA) and LMB (Hamamoto et al., 1983a,b). To confirm the identification of the purified B21P2 ion at m/z=563.331 [M+Na]⁺ as LMB, a sample of commercial LMB (Sigma L2913-5UG) was subjected to HPLC, where it eluted at 15.2 min, and mass spectrometry, where it gave an ion at m/z=563.333 [M+Na]⁺ (Fig. 3C). Additionally, the fragmentation pattern of the LMB-derived ion at m/z=563.331
[M+Na]+ (Fig. 3E) was identical to the fragmentation pattern of the B21P2-derived ion at \( m/z = 563.333 \) [M+Na]+ (Fig. 3D) confirming that our initial identification of the B21P2 bioactive compound as LMB was correct.

**Elongated cell phenotype induced by T342**

Next, we investigated activities causing an elongated cell phenotype (Fig. 1I): this phenotype is characteristic of a block in cell cycle progression while cells continue to grow (Bonatti et al., 1972) and screening for this phenotype has been instrumental in identifying many of the genes required for cell cycle progression (Nurse et al., 1976; Nurse and Thuriaux, 1980; Nasmyth and Nurse, 1981). One way in which the cell cycle could be blocked is by activation of a checkpoint delaying entry into mitosis in response to DNA damage or stalled replication forks (Enoch et al., 1992). Inability to activate the DNA checkpoint in the presence of DNA damage or stalled replication forks allows cells to enter mitosis to give a cut phenotype (Kelly et al., 1993; Saka and Yanagida, 1993). Cds1 is required to activate the intra-S checkpoint when DNA replication is stalled and Chk1 is activated in response to DNA damage. Rad3 is an upstream activator of both pathways (al-Khodairy et al., 1994). Actinomycetes are known to produce DNA-damaging compounds, for example, the anti-cancer agents bleomycin (Umenzawa et al., 1966) and adriamycin (Arcamone et al., 1969). We decided to focus on T342 as a representative example of an elongation-inducing strain and to investigate whether this phenotype was due to checkpoint activation by the bioactive natural product compound or a block in cell cycle progression in the absence of checkpoint activation.

**T342-induced cell elongation is Chk1 dependent**

When T342 extract was added to a Rad52–GFP-expressing strain (Table S2), the number of Rad52 foci increased compared to that found in cells in the absence of T342 suggesting that cell elongation in the presence of T342 is due to checkpoint activation as a result of either stalled DNA replication or DNA damage (Lisby et al., 2001) (Fig. S1). To test these two possibilities, we added semi-purified T342 extract to 972h- cells (T342 is active in both wild-type 972h- and SAK950 cells to a similar level; see Fig. S2, compare A with C), and rad3Δ, chk1Δ and cds1Δ strains (Table S2). Wild-type 972h-cells and cells lacking Cds1 elongated, suggesting that Cds1 (required for the intra-S checkpoint) is not required for checkpoint activation in the presence of T342 (Fig. 4A,B,E,F). Cells lacking Chk1 (required for the DNA damage checkpoint) or Rad3 (required upstream of both Cds1 and Chk1) did not elongate but entered mitosis prematurely to give a cut phenotype (Kelly et al., 1993; Saka and Yanagida, 1993; Yanagida, 1998) (Fig. 4B,C,G,H). These results demonstrate that the elongation phenotype was due to activation of the Chk1-dependent DNA checkpoint and suggest that T342 causes DNA damage.
Fig. 4. T342 activates the DNA damage pathway. (A) Wild-type 972h- cells in the absence of T342 or (B) 6 h after addition of T342. (C) rad3Δ cells in the absence of T342 or (D) 6 h after addition of T342. (E) cds1Δ cells in the absence of T342 or (F) 6 h after addition of T342. (G) chk1Δ cells in the absence of T342 or (H) 6 h after addition of T342. Cells were stained with DAPI to show DNA, and the white arrows show cells with a cut phenotype. Both 972 h- and cds1Δ cells elongated in the presence of T342; no cut cells were observed (n=200 cells). Neither rad3Δ nor chk1Δ cells showed any elongation (n=200 cells). For chk1Δ, 32% of cells and for rad3Δ 30% of cells showed abnormal DNA segregation (cut phenotype) (n=110 cells and n=100 cells, respectively). Scale bars: 10 µm.

Swollen cell tip phenotype induced by 5307(1B)
The phenotype induced by 5307(1B) was initially observed during the screen as a thickened region, mainly at one of the cell tips (Fig. 1C). Using a purer, more concentrated, extract we showed that the phenotype was due to accumulation of cell wall or septal material (Calcofluor staining material) at one of the cell tips (Fig. 6A,B) causing a swollen cell tip. At late time points, the cell tip became dramatically swollen (data not shown).

We purified the natural product produced by 5307(1B) that was responsible for this phenotype to conclusively identify the activity. We found that medium A was the most effective production medium for the bioactive compound after 7 days of growth. Using these conditions, agar plates were harvested and the bioactive compound was extracted by reverse-phase flash chromatography followed by preparative HPLC and mass spectrometry as described in the Materials and Methods. Despite not being detected during the HPLC using any of the wavelengths used by the diode array detector (DAD) array, two fractions eluting at between 10 and 10.5 min containing the active compound was identified by filter disc bioassay. These were subjected to mass spectrometry and both revealed the presence of an ion at m/z=529.135 [M+Na]⁺ (Fig. 5A). Querying of the Dictionary of Natural Products (DNP) for a mass of 529.135 revealed that our initial identification was correct.

Identification of the bioactive compound, which eluted in the HPLC gradient at 11.5 min and produced an ion at m/z=529.135 [M+Na]⁺ (Fig. 5A). Querying of the Dictionary of Natural Products suggested that the most likely candidate was streptonigrin (SN), which possesses a very similar mass to that reported here. To confirm the identity of the purified T342 activity as SN, a sample of commercially purified SN (Sigma) was subjected to HPLC, where it eluted at 11.5 min, and mass spectrometry, where it gave an ion at m/z=529.135 [M+Na]⁺ (Fig. 5C). The fragmentation pattern of the ion at the m/z=529.1330 species [M+Na]⁺ generated by SN (Fig. 5D) was identical to the fragmentation pattern of the ion at m/z=529.1338 species [M+Na]⁺ from T342 (Fig. 5B) confirming that our initial identification was correct.

Additionally, we tested commercially available SN in a S. pombe filter paper assay to see whether this gave a similar phenotype to the purified bioactive compound from T342. We found that the phenotype using SN was indistinguishable from that seen with T342 (Fig. S2, compare C with D). Our results suggest that SN leads directly or indirectly to DNA damage, as previously reported (see Discussion).

Purification and identification of the bioactive compound from T342
Medium H (GYM) agar plates were inoculated with T342 and grown for 7 days, after which the plates were harvested and the bioactive compound was extracted in a multi-step procedure as described above. This initially comprised a ‘freebasing’ process followed by reverse-phase flash chromatography, preparative HPLC and mass spectrometry. The purification process resulted in identification of a bioactive compound, which was extracted by reverse-phase flash chromatography, preparative HPLC and mass spectrometry. The purification process resulted in...
higher concentration of commercial cycloheximide (10 mg ml\(^{-1}\)) was used with wild-type cells, SAK950 cells died and did not exhibit the cell wall phenotype.

**Rounded cell phenotype induced by IS1**

The final phenotype we investigated was the rounded cell phenotype (Fig. 1G) in the presence of IS1. This rounded category included phenotypes that varied from round cells to wider cells that were still rod-shaped. IS1 produced a mixed phenotype of rounded cells and more rod-shaped cells (Fig. 8C,D). Previous work has shown that rounded/wide cells can be the result of a number of different cellular defects. For example, defects in nutrient monitoring, growth zone size and assembly, actin delocalisation or cell wall defects can all generate this phenotype (Ribas et al., 1991; Mendoza et al., 2005; Leonhard and Nurse, 2005; Kelly and Nurse, 2011; Thompson and Sahai, 2015; Verde et al., 1995; Kume et al., 2013).

The compound produced by IS1, which was responsible for inducing the rounded cell phenotype, was successfully identified using the protocol described in the Materials and Methods. A peak common to the active fractions was identified, and a highly purified extract of the activity, which possesses a distinctive triple-peak absorbance pattern (Fig. 8A), was obtained by ‘peak picking’. A sample of this compound was subjected to mass spectrometry and revealed the presence of an ion at \(m/z=693.379\) [M+H]\(^{+}\) (Fig. 8B). Querying of the Dictionary of Natural Products for an accurate mass of 692.371 did not produce any matches indicating, *prima facie*, that the compound represents a novel chemical entity.

To confirm that the activity purified from IS1 was responsible for the rounded cell phenotype, we carried out a filter paper assay against SAK950 cells and showed that a novel polyene induced the rounded cell phenotype (Fig. 8C,D). We identified IS1 as a polyene on the basis of the triple-peak absorbance pattern in the region of \(\lambda=300–450\) nm, which is characteristic of polyenes and is used as a diagnostic aid to identify polyenes during natural product screening, see the Discussion for further details.

**DISCUSSION**

In this study, we have screened 657 actinomyces strains for secretion of activities that affect the cell shape of fission yeast. From the set of 420 strains that produced bioactivities activities, there were 149 extract of the activity, which possesses a distinctive triple-peak absorbance pattern (Fig. 8A), was obtained by ‘peak picking’. A sample of this compound was subjected to mass spectrometry and revealed the presence of an ion at \(m/z=693.379\) [M+H]\(^{+}\) (Fig. 8B). Querying of the Dictionary of Natural Products for an accurate mass of 692.371 did not produce any matches indicating, *prima facie*, that the compound represents a novel chemical entity.

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**DISCUSSION**

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strains that we decided to follow up and here we have reported the identification by visual screening, and the purification and preliminary analysis, of four of these bioactivities as a proof of principle for this type of approach.

**Leptomycin origin and asymmetric phenotype bioactivity**

The leptomycins were initially identified in 1983 (Hamamoto et al., 1983a,b), and the effect of LMB on *S. pombe* was studied soon after their purification (Hamamoto et al., 1985). It was shown to cause cell elongation, and an inhibitory effect of a high concentration of LMB (10 µg ml$^{-1}$) on DNA synthesis was observed, although the primary effect was suggested to be inhibition of mitotic exit without blocking septation (Hamamoto et al., 1985). LMB-resistant mutants in *S. pombe* resulted in the identification of *crm1* as the cellular target and that LMB binds covalently to Crm1 at a cysteine residue in a conserved region of the protein (Kudo et al., 1998; Kudo et al., 1999). Crm1 mediates the export of proteins possessing nuclear export signals (NES) from the nucleus to the cytoplasm (Kudo et al., 1998; Fukuda et al., 1997; Stade et al., 1997) and has been implicated in a number of different cellular processes in different organisms, including centromere and telomere arrangement (Funabiki et al., 1993). Crm1 is linked to the spindle pole body (SPB) by Spc72 (Neuber et al., 2008) and is involved in locating pericentrin to the centrosome (Liu et al., 2009). In addition, it is associated with nucleophosmin and is involved in regulating centrosome duplication (Wang et al., 2005; Shimamura et al., 2005). It has also been shown that centrin and pericentrin accumulate in the nucleus following treatment with LMB (Keryer et al., 2003). LMB competes with the NES for stable formation of a complex with Ran-GTP. Ran-GTP stabilises microtubule plus-ends and has been shown to concentrate at the centrosome (Keryer et al., 2003), and in *S. pombe*, it interacts with at least one component of the SPB and controls microtubule integrity (Fleg et al., 2000).

Our results suggest that a significant effect of LMB is on the microtubule cytoskeleton, and we suggest that this leads directly to mis-positioning of the nucleus and septum. Although the LMB phenotype has been well characterised (see above), this is the first time that a septum-positioning defect has been reported for LMB in *S. pombe*. It is tempting to interpret the major LMB-induced *S. pombe* cell morphology phenotype in terms of disruption of Crm1 functions relating to the export of factors affecting the interphase microtubules, but it is possible that there are other factors whose Crm1-dependent export/localisation are also affected. Indeed, it has been shown that the localisation of 285 proteins, many of which are involved in cell cycle, cell division and cell morphology processes, are dependent on Crm1, and therefore subject to LMB-mediated disruption (Matsuyama et al., 2006).

Many tumour suppressors and oncoproteins use Crm1 for their nuclear export and, as the best-characterised nuclear export inhibitor, LMB has been a candidate for anti-cancer drug development, although it failed clinical trials due to toxicity. Nonetheless, the search for natural products that possess nuclear export inhibitory activity has continued (Cautain et al., 2014; Mutka et al., 2009) and our results show that the *S. pombe* morphological effect could provide a means of screening for novel inhibitors of this class.

**SN origin and cell elongation phenotype**

SN was first isolated from *Streptomyces flocculus* (Rao and Cullen, 1960), although identical molecules have also been isolated from other species; for example, ‘bruneomycin’ from *Actinomycetes albus var bruneomycin* (Brazhnikova et al., 1968). The structure of SN has been elucidated (Rao et al., 1963) and it possesses an aminoquinone moiety, in common with many other bioactive agents of actinomycete origin, including mitomycin C, actinomycin, rifamycin and geldanamycin. SN has been shown to exert its anti-tumour effects through direct and indirectly acting mechanisms (reviewed in Bolzán and Bianchi, 2001). SN is an aminooquinone moiety, in common with many other bioactive agents of actinomycete origin, including mitomycin C, actinomycin, rifamycin and geldanamycin. SN has been shown to exert its anti-tumour effects through direct and indirectly acting mechanisms (reviewed in Bolzán and Bianchi, 2001). SN acts to damage DNA directly when it forms complexes with metal ions, including Cu$^{2+}$ (Sugiuira et al., 1984), that are capable of intercalating with DNA (Cone et al., 1976). Also, when the quinone moiety undergoes autoxidation to semiquinone, in the presence of NADH, reactive oxygen species (ROS) capable of cleaving DNA are produced (Lown et al., 1978). However, SN also...
damages DNA indirectly through inhibition of topoisomerase II activity by binding to the minor groove and stabilising the transesterification intermediate of the enzyme, so inhibiting re-ligation of the DNA strands and generating DNA breaks (Yamashita et al., 1990; Capranico et al., 1994). It has also been shown through in vitro experiments that SN inhibits the ATPase activity of *S. cerevisiae* Rad54 by generating ROS and so prevents Holliday junction branch migration (Deakayne et al., 2013).

In vivo, SN works through both the direct and indirect DNA-damaging mechanisms mentioned above, depending on its concentration, with the inhibition of topoisomerase function being the predominant mechanism at lower concentrations (Legault et al., 1997).

**Cycloheximide origin and cell wall defect phenotype**

One of the more interesting aspects of the present study is the fact that in several cases the phenotypes observed during the screen are not due to a direct effect of the natural product compound on its cellular target but are due to downstream effects caused by inhibition of the primary target system. The cell wall defect phenotype observed to occur in response to 53017(1B) was initially expected to be due to an activity affecting cell wall biosynthesis, which led to a build-up of aberrant cell wall material within the cell and consequently to the observed distortions of cellular morphology. However, isolation of the active compound indicated that it was cycloheximide, whose identification was consistent with the similarity between the 16S rRNA gene sequence of strain 5307(1B) and that of *Streptomyces griseus*, which produces cycloheximide (actidione) (Whiffen et al., 1946; Leach et al., 1947; Kornfeld and Jones, 1948). Cycloheximide is well known as a general protein synthesis inhibitor (Kerridge, 1958) and is widely used as a research tool; therefore, the severe effects on the cell wall were unexpected. However, both cycloheximide and 5307(1B) extract induce a similar phenotype, with the *S. pombe* cell wall becoming thickened mainly at one of the cell tips. To our knowledge, this is the first time that an effect of cycloheximide on *S. pombe* cell morphology has been reported.

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**Fig. 8.** The purified novel compound derived from IS1 causes a rounded cell phenotype. (A) UV and visible absorbance spectrum of the IS1 compound. (B) Mass spectrometry data for compound purified from IS1 [M+H]+. (C) SAK950 control (left panel), SAK950 cells from the plug assay with IS1 (middle panel), and SAK950 cells from filter disc assays with ‘peak-picked’ purified IS1 compound (right panel). (D) Representative cells of SAK950 control (left panels) and SAK950+peak picked IS1 from B shown at higher magnification (right panels). Scale bars: 10 µm.
The differential effects of cycloheximide on the synthesis of the cell wall components mannan and glucan in *S. cerevisiae* have been noted previously, with mannan synthesis being inhibited, whereas glucan synthesis remains unaffected for a considerable time post cycloheximide treatment (Elorza et al., 1976; Sentandreu and Lampen, 1970; Coen et al., 1994). The thickening of the *S. cerevisiae* cell wall following cycloheximide treatment has also been noted, particularly in the septum region at the tip of the bud (Rodriguez et al., 1979). Although the effect of cycloheximide treatment on *S. pombe* division has been investigated and the presence of thickened cell plates observed in dividing cells (Polanshek, 1977), the more-severe phenotype observed in this study has not been reported previously. Cycloheximide inhibition of protein synthesis in *S. pombe* may, as in *S. cerevisiae*, have a differential effect on mannan and glucan synthases. If so, then lack of an effect of cycloheximide on glucan synthases for a period of time after addition is likely to lead to the accumulation of unincorporated glucan at the septum in dividing cells and at the new cell tips generated at cell division, producing the aberrant cell wall *S. pombe* phenotype that we observed.

**Identification of the novel IS1 compound**

The novel compound IS1 was identified through its ion at *m/z*=693.379 [M+H]+ by a search of the Dictionary of Natural Products. The determination of the absorbance spectrum of the molecules (Fig. 8A) indicated that it possesses a triple-peak pattern that is characteristic of molecules possessing a plurality of conjugated unsaturated carbon–carbon bonds (Oroshnik et al., 1955). Antifungal compounds that possess similar triple-peak absorbance spectra include the polyene macrolides (Oroshnik et al., 1955; Hamilton-Miller, 1973), and the triple-peak absorbance spectrum is widely used as a diagnostic tool in natural product screening to identify polyenes (Lemris et al., 2003; Ouchdouch et al., 2001). Based on the maxima of the absorption spectrum, we would predict that five or six unsaturated bonds are present (Hamilton-Miller, 1973) and, as during analysis of the IS1 compound by LC-MS a series of ions differing by 18 mass units were generated (data not shown), it is likely the IS1 compound possesses a multiplicity of hydroxyl groups that are lost via water as the major ion species fragments.

Additionally, the cell phenotype induced by IS1 is similar to but not identical to the rounded phenotype induced by the polyene antifungal nystatin (data not shown) and may reflect a similar mode of action. Sterols are well known to play an important role in maintaining the integrity of lipid rafts, which are responsible, at least in part, for the correct localisation of protein complexes necessary for cell morphology/polarity (Takeda et al., 2004; Takeda and Chang, 2005; Wachtler et al., 2003; Codlin et al., 2008; Fischer et al., 2008). Although we do not attempt a precise explanation of the cell phenotype in terms of the action of the novel IS1 compound, or indeed of polyenes in general, we note that the rounded/wide phenotypes are consistent with reduced control of cell morphology/polarity which leads to loss of, or a less defined, rod-shaped phenotype.

The drug-sensitive strain used in this study, SAK950, possesses an *erg5A* mutation (Aoi et al., 2014) and so is deficient in ergosterol (Iwaki et al., 2008), which is known to be a target molecule of polyenes (Gray et al., 2012). Interestingly, although it was expected that SAK950 would therefore show some resistance to polyenes its sensitivity to nystatin is almost identical to the wild-type strain 972h-, although it is more sensitive than wild-type cells to other available antifungal agents (data not shown). Therefore, it seems that polyenes, in addition to targeting ergosterol itself, may also be able to exert a killing effect by interacting with the ergosterol precursors present in SAK950.

**Conclusions**

Recently there has been a call for “more concerted screening campaigns of actinobacterial metabolites against model eukaryotes… These screens should harness more than just live/dead screening. In other words, we should look for interesting developmental and behavioural phenotypes using well-developed model systems” (Ho and Nodwell, 2016). The screen described here aptly fits the above description; we have successfully screened strains producing putative anti-fungal natural products, from a major collection of actinomycete bacteria, for their ability to induce cell shape defects in *S. pombe*, with the aim of identifying chemical entities capable of exerting effects on the cell cycle/cell morphology. We were successful in identifying many such strains, which induced a wide range of aberrant phenotypes on *S. pombe*. The phenotypic groups we observed are similar to, but not completely overlapping with those observed after screening of the *S. pombe* genome-wide gene deletion collection (Hayles et al., 2013). As expected, the complexity of phenotypes we have identified suggests that bioactive compounds may induce a different range of phenotypic responses than a single gene knockout would generate, so demonstrating the richness and complexity of the biological activities the present study has identified.

Following on from the initial screen, we identified and purified four compounds, LMB, SN and cycloheximide produced by actinomycete bacteria B21P2, T342 and 5307(1B), respectively, and a novel compound from actinomycete bacterium IS1. These results demonstrate that this type of screen will identify both known compounds of biological interest and novel compounds. We have analysed the *S. pombe* phenotypes in the presence of these drugs and found that they correlate with the known or predicted bioactivities of these compounds. These four case studies confirm the validity of our approach to identify bioactivities affecting *S. pombe* cell cycle and cell morphology, and suggest that some of the other 149 activities that we are currently investigating will be novel, biologically important activities affecting these processes in *S. pombe*. Given the conservation of genes encoding proteins involved in the cell cycle and microtubule/actin cytoskeleton between *S. pombe* and higher eukaryotes, and the medical use of some of the compounds affecting the *S. pombe* cell wall, which may act as antifungals, it is likely that novel natural products identified in this way will also be relevant to higher eukaryotes.

**MATERIALS AND METHODS**

**Yeast strains, growth conditions and media**

Strains used in this study (Table S2) were grown at 30°C in YE4S (Moreno et al., 1991) unless otherwise stated.

**S. pombe morphological assays**

**Plug assay**

1 ml pipette tips were used to cut agar plugs from lawns of bacterial strains on GYM agar plates (see below). A plug from each of three different bacterial strains plus an agar-only control plug was then placed in a quadrant on a YE4S plate seeded with 5×10^5 mid-log phase SAK950 cells. Plates were incubated overnight at 30°C and screened the following day for the presence or absence of a halo in the *S. pombe* lawn surrounding the plugs and microscopically for the phenotype of the *S. pombe* cells (Fig. 1). This procedure was carried out after 5, 7 and 9 days of bacterial growth. As biomolecules diffuse through the agar at different rates and may have...
different effects at different concentrations, the phenotype of the cells was examined from the plug to the edge of the halo to ensure all phenotypes were observed (Table S1).

Filter disc assay
Filter discs soaked in extract (10–30 μl with an appropriate amount of H2O or solvent) (Whatman cat no. 2017-006) instead of bacterial plugs were used as described above.

DNA damage and checkpoint activation
To assess DNA damage mediated by T342, the T342 extract was added to an exponentially growing culture of rad52GFP (Lisby et al., 2001) in YE4S liquid medium and the number of Rad52-GFP foci at 0, 1, 3 and 5 h after addition were counted and compared to the number upon no T342 extract addition.

To assess checkpoint activation mediated by T342, semi-purified T342 extract (1:100 dilution) was added to 972h−, rad3Δ, cdc15Δ and chk1Δ S. pombe strains growing exponentially in YE4S medium. After 6 h, the cell phenotype was observed to see whether cells had elongated or had a cut phenotype (shown by DAPI staining).

Microscopy
Bright-field images of live cells and DAPI-stained cell images were acquired on a Zeiss Axioskop 40 microscope with a Zeiss 63×/1.4 NA oil ph3 objective and a Zeiss Axioscan MRm camera. For the plug assay plates, cell images were captured with a Sony HD AVCHD camera using a Zeiss Axioskop 40 microscope with a 50×/0.56 NA Nikon objective and 2.5× Optovar magnification changer. PN5352 and Rad52–GFP live cells were imaged using a DeltaVision microscope (Applied Precision) and a 60×/1.42 NA oil immersion lens. A total of ten z-sections with 0.4 μm spacing were taken with a Coolscope-HQ digital CCD camera (Roper Scientific) and deconvolved using SoftWoRx (Applied Precision). For time-lapse imaging, cells were grown in YE4S medium at 32°C to early exponential growth, B21P2 was added and cells were incubated for a further 40 min before transferring to the microfluoride cell culture system (CellASIC, ONIX). Six z-sections with 0.6 μm spacing were acquired for each time point with 5 min time intervals for 245 min at 32°C using the DeltaVision microscope (Applied Precision). For the SAK950 cells shown in Fig. 8D, live cells were imaged using a DeltaVision microscope (Applied Precision) with a 60×/1.42 NA oil immersion lens after resuspending cells from the edge of colonies in water.

Reagents were used at the following concentrations/dilutions: purified B21P2 at 1:100 dilution, LMB (Sigma L2913-5UG) at 1 mg/ml, Calcofluor (MD Biomedical) at 1:100 dilution of water for 10 min and Solid media had agar added (10 g l−1). Strain B21P2 was isolated from the rhizosphere of Paraseriannes falcatoria (Sembringer, 2000); the most closely related species is Kitasatospora pufferickiae (16S RNA gene has 8/1386 bp mismatches).

Strain T342 was first described in Sahin, 1995. The most closely related species is Streptomyces albus (16SrRNA gene has 14/1438 bp mismatches). The most closely related species to strain 5307(1B) (ATCC 12433) is Streptomyces griseus (16SrRNA gene has 0/1379 bp mismatches). The most closely related species to strain IS1 is Streptomyces misionensis (16S RNA gene has 4/1414 bp mismatches).

Production of water- and acetone-based bacterial cell extracts
B21P2 and IS1 were streaked onto GYM agar plates and grown for 7 days. Production of the bioactivity was assessed by using the plug assay. Agar from the plates was harvested, disrupted by passing through a 50 ml syringe and frozen overnight at −20°C. A water-based extract was produced by squeezing the thawed slurry through muslin cheesecloth and filtering the resultant liquid through Whatman filter paper. A concentrated extract was obtained by freeze drying the filtered extract in an Edwards ‘Modulyo’ freeze dryer and resuspending the lyophilised material in an appropriate volume of water. The agar–cell mixture produced was resuspended in a volume of acetone equal to the initial volume of water-based extract and incubated overnight at 4°C. Acetone-based extract was produced by squeezing the mixture through muslin cheesecloth, filtering it and removal of acetone using a Bühci vacuum rotary evaporator to generate a concentrated extract comprising the residual water present in the spent agar–cell slurry.

Production of bioactive crude extracts
B21P2 was streaked onto INAS agar plates, T342 onto GYM agar plates and 5307(1B) onto Medium A plates. All strains were grown for 7 days and production of bioactive compounds assessed by use of the plug assay. Agar from the plates was harvested, disrupted by passing through a 50 ml syringe and frozen overnight at −20°C. A water-based extract was produced by squeezing the thawed slurry through muslin cheesecloth and filtering the resultant liquid through Whatman filter paper. Washed Amberlite™ XAD16N resin beads (Sigma) were added (20 g l−1) to the resultant liquid to adsorb bioactive compounds. The liquid was successively extracted three times with portions of Amberlite™ resin beads, and the compounds released by washing the resin with volumes of methanol equal to the volume of the liquid extract. The methanol was removed by rotary evaporation (Büchi) to obtain water-based concentrates.

Reverse-phase flash chromatography
The B21P2- and 5307(1B)-derived water-based concentrates, and IS1-derived acetone-based concentrates were passed through a pre-packed disposable reverse-phase C18 25 g SNAP flash chromatography column (Biotage) and the bound compounds eluted in a 0–100% water (0.1% formic acid)–methanol (0.1% formic acid) gradient using a Biotage Isolera One chromatography machine. T342-derived extract produced by resuspension of ‘freebase’ material (see ‘Solvent Extraction’ below) in water was treated similarly. For all strains, fractions containing bioactive compounds were identified by spotting 30 μl aliquots onto filter paper discs, which were used in the filter disc assay. Active fractions producing a halo of cells with the expected characteristic phenotypes were pooled and the methanol removed by a GeneVac Series II system equipped with a GeneVac VC3000TA condenser unit.

Thin layer chromatography
B21P2- and 5307(1B)-derived samples were subjected to thin layer chromatography (TLC) using silica gel plates (Fluka Analytical) and a range of solvents: ethyl acetate, methanol and diethyl ether. After running the samples, the TLC plates were pressed, silica side down, onto YE4S agar plates and the position of the TLC plates, sample loading site and solvent front were recorded on the back of the petri dish. After a 30 min transfer period the TLC plates were removed and the YE4S agar plates were inoculated with S. pombe in a similar fashion to in the plug and filter paper assays. After an overnight incubation at 30°C, zones of inhibition in the S. pombe lawns comprising cells of aberrant morphology indicated the presence of bioactive compounds of interest and comparison of the location of these zones with the solvent front and sample loading site markings allowed assessment of the solubility of the bioactive compound in the
various solvents and selection of the most appropriate solvent(s) for future purification steps.

Solvent extraction
Ethyl acetate extractions of B21P2- and 5307(1B)-derived material were obtained by shaking the aqueous extract with an equal volume of ethyl acetate in a separating funnel for 5 min followed by separation of the two phases. The ethyl acetate phase was rotary evaporated with silica so as to adsorb the compounds therein. A ‘freebas ing’ process was employed with regard to T342-derived extracts where the pH of the aqueous extract (pH 8) was raised to pH 10 and an ethyl acetate extraction used to remove contaminants in the organic phase. This was followed by alteration of the pH of the aqueous phase to pH 4 and again obtaining an ethyl acetate extraction where the bioactive compound migrated into the organic phase, after which it was recovered by rotary evaporation (Buchi) and resuspended in water.

Normal-phase flash chromatography and size exclusion chromatography
The silica with adsorbed B21P2-derived compound was transferred into a pre-packed, disposable K-Sil 25 g SNAP column (Biotage) and used in normal-phase flash chromatography with elution in a 0–100% diethyl ether–ethyl acetate gradient. The silica with adsorbed 5307(1B)-derived compound was similarly treated with elution in a 0–100% dichloromethane–methanol gradient. For both strains, fractions containing bioactive compounds were identified by spotting 30-µl aliquots onto filter paper discs which were used, as described above, to identify active fractions producing a halo comprising cells exhibiting the expected characteristic phenotypes. For B21P2, active fractions were pooled, the solvent removed using a GeneVac Series II (Holloway) and the resultant sample was loaded onto a 150 PSI Max (Omnifit) size exclusion column packed with Sephadex™ LH-20 connected to a Biotage Isolera One chromatography machine and eluted in a methanol mobile phase. Active fractions, identified by the filter paper assay, were pooled and the methanol removed using a GeneVac Series II.

HPLC and mass spectrometry
The B21P2-derived compound was solubilised by addition of 25% methanol, and 100 µl aliquots were passed through a Perkin Elmer Series 200 HPLC machine equipped with a Phenomenex 150 × 4.6 mm Synergy 4 µm reverse-phase column and a Hichrom C18 guard column and eluted using a 43 min 25%–100% water (0.1% formic acid)–acetonitrile (0.1% formic acid) gradient, with a flow rate of 1 ml min⁻¹ and monitoring at λ=254 nm. Fractions (0.5 ml) were collected in a 96-well block, and active fractions identified by the filter paper assay. Following correlation of peaks in the UV absorbance trace with the active fractions, the method was repeated with an Agilent Technologies 1260 Infinity liquid chromatography machine equipped with an integrated fraction collector so as to ‘peak pick’ and collect column eluate corresponding to the peaks of interest. The gradient used was 25%–100% water (0.1% formic acid)–acetonitrile (0.1% formic acid) for 15 min ending with a 5 min 100% acetonitrile column wash using a flow rate of 1 ml min⁻¹ and monitoring with a DAD array at λ=210, 254, 273, 280, 300, 350 and 400 nm relative to λ=360 nm. T342- and 5307(1B)-derived compounds were similarly analysed but were eluted in a gradient of 15%–100% water (0.1% formic acid)–acetonitrile (0.1% formic acid). ISI-derived compound was passed through an Agilent 150 × 4.6 mm Eclipse Plus C18 3.5 µm reverse-phase column and a Hichrom C18 guard column and eluted using a 30 min 30%–100% water–acetonitrile gradient, with a flow rate of 1 ml min⁻¹ and monitoring at λ=350 nm with an Agilent Technologies 1260 Infinity liquid chromatography machine equipped with an integrated fraction collector so as to peak pick and collect column eluate corresponding to the peak of interest.

After removal of solvent, by using the GeneVac Series II, and addition of 25% methanol to enhance solubility, the samples were analysed by electrospray mass spectrometry (ESI-MS) using a LTQ-FT (Thermo) mass spectrometer with a 7T magnet at the Pinnacle Laboratory (University of Newcastle). Experiments were run with a parent/precursor scan at 100,000 resolution. MS/MS fragmentations were carried out in the ion-trap (LTQ) stage of the instrument. Commercially available LMB (Sigma), SN (Sigma) and cycloheximide (Melford Laboratories) were used as comparative standards for the purified molecules of interest. For HPLC, LMB (6.5 µg/ml) and SN (20 µg/ml) were made up in 20% methanol, and for mass spectrometry, LMB (11 µg/ml) was in 50% methanol whereas SN (20 µg ml⁻¹) and cycloheximide (10 µg ml⁻¹) were in 20% methanol.

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Competing interests
N.E.E.A. and J.E. hold shares, or share options, in Demiris Ltd.

Author contributions
Conceptualization: R.A.L., J.H., P.M.N.; Methodology: R.A.L.; Formal analysis: R.A.L., J.L.; J.H.; Resources: N.E.E.A., J.E., P.M.N.; Writing - original draft: R.A.L., J.H.; Writing - review & editing: R.A.L., J.H., P.M.N.; Project administration: J.E.; Funding acquisition: P.M.N.

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Supplementary information
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