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Identification of the First ATRIP–Deficient Patient and Novel Mutations in ATR Define a Clinical Spectrum for ATR–ATRIP Seckel Syndrome

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Abstract

A homozygous mutational change in the Atraxia-Telangiectasia and RAD3 related (ATR) gene was previously reported in two related families displaying Seckel Syndrome (SS). Here, we provide the first identification of a Seckel Syndrome patient with mutations in ATRIP, the gene encoding ATR–Interacting Protein (ATRIP), the partner protein of ATR required for ATR stability and recruitment to the site of DNA damage. The patient has compound heterozygous mutations in ATRIP resulting in reduced ATRIP and ATR expression. A nonsense mutational change in one ATRIP allele results in a C-terminal truncated protein, which impairs ATR–ATRIP interaction; the other allele is abnormally spliced. We additionally describe two further unrelated patients native to the UK with the same novel, heterozygous mutations in ATR, which cause dramatically reduced ATR expression. All patient-derived cells showed defective DNA damage responses that can be attributed to impaired ATR–ATRIP function. Seckel Syndrome is characterised by microcephaly and growth delay, features also displayed by several related disorders including Majewski (microcephalic) osteodysplastic primordial dwarfism (MOPD) type II and Meier-Gorlin Syndrome (MGS). The identification of an ATRIP–deficient patient provides a novel genetic defect for Seckel Syndrome. Coupled with the identification of further ATR–deficient patients, our findings allow a spectrum of clinical features that can be ascribed to the ATR–ATRIP deficient sub-class of Seckel Syndrome. ATR–ATRIP patients are characterised by extremely severe microcephaly and growth delay, microtia (small ears), micrognathia (small and receding chin), and dental crowding. While aberrant bone development was mild in the original ATR–SS patient, some of the patients described here display skeletal abnormalities including, in one patient, small patellae, a feature characteristically observed in Meier-Gorlin Syndrome. Collectively, our analysis exposes an overlapping clinical manifestation between the disorders but allows an expanded spectrum of clinical features for ATR–ATRIP Seckel Syndrome to be defined.

Introduction

Seckel Syndrome (SS) (OMIM 216000) is an autosomal recessive disorder characterised by marked microcephaly, intra-uterine and post-natal growth retardation, developmental delay and characteristic facial features, encompassing micrognathia (small and receding chin), receding forehead and pronounced nose [1]. Majewski (microcephalic) osteodysplastic primordial dwarfism (MOPD) type II and Meier-Gorlin Syndrome (MGS) also display microcephaly and primordial dwarfism [2,3]. How-
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Author Summary

Seckel Syndrome (SS) is a rare human disorder characterised by small head circumference and delayed growth. Patients can show additional features including abnormal bone development, receding chins, sloping foreheads, and small ears. In 2003, we identified ataxia telangiectasia and Rad3 related (ATR) as a causal genetic defect in two related families displaying SS. However, additional patients with mutations in ATR have not hitherto been identified. Here, we describe two further patients with novel mutations in ATR. Additionally, we identify a patient with mutations in ATRIP, which encodes an interacting partner of ATR, representing a novel genetic defect causing SS. ATR functions to promote the ability of cells to recover from difficulties encountered during replication. We show that patient-derived cells have reduced ATR and ATRIP protein levels and defective ATR/ATRIP function. Our identification of further ATR–ATRIP defective patients and a characterisation of their clinical features aids the characterisation and identification of this form of SS and provides insight into the role played by the ATR–ATRIP complex during development.

ever, each of these disorders display an additional spectrum of features conferring clinical distinction. Despite this, on an individual basis, assigning patients to a specific classification is difficult. Additionally, primary microcephaly represents a disorder displaying pronounced microcephaly without marked impact on growth [4]. Five loci conferring SS have been described with four genes identified [5,6]. The first causal genetic defect identified for SS was the Ataxia-Telangiectasia and Rad3 related (ATR) gene [7]. A homozygous mutation in ATR was identified in two related SS families and cell-based studies provided strong evidence for an impact on ATR function in patient cell lines. This sub-class of SS was designated ATR–SS. More recently, mutations in CTIP were identified in a SS patient as well as in a family described as displaying Jawad Syndrome [8]. Additionally, mutations in CENPJ and CEP152, two centrosomal proteins, have been described in SS patients, although mutations in these genes more frequently confer primary microcephaly [9,10]. Mutations in PCNT, which encodes a centrosomal protein, and ORC1L, a component of the original licensing complex, were reported in patients originally classified as SS although in both cases retrospective analysis revealed that such mutations more frequently cause MOPD type II or MGS, respectively, highlighting the diagnostic challenge faced in the clinic [11–15]. These studies demonstrate that evaluation of multiple patients is required to provide insight into the spectrum of clinical features conferred by specific gene defects, which ultimately aids an understanding of the role of the defective protein during development. To date all ATR–SS patients belong to one of two related families, which harbour the identical homozygous mutation in ATR, thereby limiting the characterisation of the clinical phenotype conferred by ATR deficiency. Furthermore, no patients deficient in ATR interacting protein, ATRIP, which is required for ATR stability, have hitherto been described.

ATR, like the related Ataxia-Telangiectasia mutated (ATM) protein, is a phosphoinositide-3 kinase (PI3)-like kinase that functions at the centre of a signal transduction network activated by DNA damage, and most importantly, by replication fork stalling [16]. ATR and ATM share phosphorylation targets but whilst ATM is activated by DNA double strand breaks (DSBs) that arise, for example, following exposure to ionising radiation (IR), ATR is activated by single stranded (ss) regions of DNA that arise following replication fork stalling or exposure to agents that induce bulky DNA adducts [17,18]. Importantly, since replication fork stalling occurs during most cycles of replication, ATR is essential. ATM, in contrast, is non-essential presumably because endogenous DSBs arise infrequently. ATR forms a stable complex with ATR-interacting protein (ATRIP), which is required for ATR stability [19]. Further, ATRIP is required for ATR localisation to ssDNA regions and hence for ATR activation. Consequently, in a range of organisms loss of ATRIP or its homologue, phenocopies ATR deficiency [17,20–22]. Although ATM and ATR share overlapping substrates, ATR specifically phosphorylates Chk1 whilst ATM phosphorylates the related kinase, Chk2. The major functions of ATR are to activate cell cycle checkpoint arrest, stabilise stalled replication forks and promote replication fork restart, which is achieved through its ability to phosphorylate a range of substrates including p53 and H2AX [18,23,24]. Interestingly, in the context of SS, ChIP promotes DNA end resection, which leads to ss DNA formation, the lesion activating ATR. Hence, ChIP functions in a mechanism leading to ATR activation. It is noteworthy that cells derived from PCNT-mutated MOPD type II patients are also defective in ATR–dependent G2/M checkpoint arrest although upstream steps in the ATR–signalling pathway are activated normally [11]. These findings suggest that PCNT is required for an important end-point of ATR function. Additionally, the origin licensing complex, components of which are mutated in MGS, is required for the initiation of replication and ORC1L-deficient MGS cell lines display slow S phase progression [13]. Similarly, ATR promotes S phase progression by facilitating recovery from replication fork stalling.

Our identification of a SS patient as well as a family described as Jawad Syndrome [8]. Additionally, mutations in CENPJ and CEP152, two centrosomal proteins, have been identified in SS patients, although mutations in these genes more frequently confer primary microcephaly [9,10]. Mutations in PCNT, which encodes a centrosomal protein, and ORC1L, a component of the original licensing complex, were reported in patients originally classified as SS although in both cases retrospective analysis revealed that such mutations more frequently cause MOPD type II or MGS, respectively, highlighting the diagnostic challenge faced in the clinic [11–15]. These studies demonstrate that evaluation of multiple patients is required to provide insight into the spectrum of clinical features conferred by specific gene defects, which ultimately aids an understanding of the role of the defective protein during development. To date all ATR–SS patients belong to one of two related families, which harbour the identical homozygous mutation in ATR, thereby limiting the characterisation of the clinical phenotype conferred by ATR deficiency. Furthermore, no patients deficient in ATR interacting protein, ATRIP, which is required for ATR stability, have hitherto been described.

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CV1780 and CV1783 displayed normal G2/M checkpoint arrest. We have previously observed that cells from other SS patients display defects in ATR–dependent G2/M checkpoint arrest but activate upstream steps in the ATR signalling cascade normally [25]. This is exemplified by cell lines from MOPD type II patients with mutations in PCNT, which are defective in ATR–dependent G2/M checkpoint arrest but proficient in ATR phosphorylation events [11]. Therefore, next, we examined whether CV1720 LBLs efficiently activate upstream steps in ATR signalling. Since these assays predominantly reflect the response of replicating phase cells, we first verified that CV1720 and control LBLs harbour a similar percentage of S phase cells (Figure S2). Pan-nuclear phosphorylation of H2AX (γH2AX) after replication fork stalling represents an ATR–specific damage response [24]. Strikingly, whilst exposure to 5 mM HU for 2 h resulted in an elevated number of cells staining positively for γH2AX in WT cells, this was not observed in either CV1720 or DK0064 (ATR–SS) cells (Figure 1B). We note that although previous studies have shown that ATM can be activated and phosphorylate γH2AX at DSBs arising following HU treatment in the absence of ATR due to enhanced fork collapse, this was not observed at 2 h post 5 mM HU exposure in these patient cells most likely due to residual ATR activity and/or the early times examined [26,27]. Chk1 represents an important ATR substrate required for G2/M checkpoint arrest. To examine Chk1 activation, we carried out Western Blotting using p-Chk1 antibodies. Following the same UV exposure conditions (2 h post 5J m\(^2\)) employed to examine G2/M checkpoint arrest, we

Table 1. Clinical features of ATR/ATRIP–deficient patients.

<table>
<thead>
<tr>
<th></th>
<th>ATRIP–SS</th>
<th>ATR–SS</th>
<th>27-4BI</th>
<th>19-8BI</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ethnicity</strong></td>
<td>Gujarati-Indian (consanguineous)</td>
<td>Pakistani (consanguineous)</td>
<td>English</td>
<td>English</td>
</tr>
<tr>
<td><strong>Birth.</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OFC (cm)</td>
<td>27.1</td>
<td>24 (-8SD)</td>
<td>27</td>
<td>24.2</td>
</tr>
<tr>
<td>Wgt (Kg)</td>
<td>2.06</td>
<td>1.1 (-3SD)</td>
<td>1.15</td>
<td>0.77</td>
</tr>
<tr>
<td>Hgt (cm)</td>
<td>NR</td>
<td>NR</td>
<td>36</td>
<td>NR</td>
</tr>
<tr>
<td><strong>Age.</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OFC (cm)</td>
<td>-9SD</td>
<td>-10SD</td>
<td>-12SD</td>
<td>-10SD</td>
</tr>
<tr>
<td>Wgt (Kg)</td>
<td>-5SD</td>
<td>-6SD</td>
<td>-3.3SD</td>
<td>-8SD</td>
</tr>
<tr>
<td>Hgt (cm)</td>
<td>-5SD</td>
<td>-6.5SD</td>
<td>NR</td>
<td>-8SD</td>
</tr>
<tr>
<td><strong>Face</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Teeth</strong></td>
<td>Dental crowding.</td>
<td>Dental crowding and malocclusion.</td>
<td>4 teeth at 20 months.</td>
<td>Dental crowding.</td>
</tr>
<tr>
<td><strong>Ears</strong></td>
<td>Small lobes.</td>
<td>Posteriory rotated with absent lobes.</td>
<td>Small, round, low set with poorly formed antihelix tragus &amp; antitragus. Absent lobes.</td>
<td>Small ears with absent lobes</td>
</tr>
<tr>
<td><strong>Hands</strong></td>
<td>Bilateral 5(^{th}) finger clinodactyly.</td>
<td>Multiple ivory epiphysis.</td>
<td>Small, tapering fingers.</td>
<td>Bilateral 5(^{th}) finger clinodactyly. 5(^{th}) metacarpels appear short. Blue colouration to both thenar eminence.</td>
</tr>
<tr>
<td><strong>Endo-crinology</strong></td>
<td>Normal IGF1, TFF, LH, FSH &amp; cortisol.</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>MRI</strong></td>
<td>14 mts: generalised cerebral atrophy, normal ventricular systems. Delayed myelination in the anterior limb of the internal capsule. Pituitary is present though of unusual shape with absent fossa.</td>
<td>NA</td>
<td>NA</td>
<td>2 yrs: abnormal gyration in posterior aspect of the cingulated gyrus extending into the parietal occipital region. Hypoplastic corpus callosum.</td>
</tr>
<tr>
<td><strong>Other</strong></td>
<td>NR</td>
<td>Developmental delay. Walked at 7 yrs.</td>
<td>No abnormal skin pigmentation. Small feet with metatarsus adductus</td>
<td>No abnormal skin pigmentation. Developmental delay. Sat at 15 mts, walked at 3 yrs 10 mts. High pitched voice, asthma, multiple chest infections, feeding difficulties-reflux (gastrostomy fed). Multiple liver cysts consistent with Caroli’s disease found at 17 mts.</td>
</tr>
</tbody>
</table>

(CV1780 and CV1783) displayed normal G2/M checkpoint arrest. We have previously observed that cells from other SS patients display defects in ATR–dependent G2/M checkpoint arrest but activate upstream steps in the ATR signalling cascade normally [25]. This is exemplified by cell lines from MOPD type II patients with mutations in PCNT, which are defective in ATR–dependent G2/M checkpoint arrest but proficient in ATR phosphorylation events [11]. Therefore, next, we examined whether CV1720 LBLs efficiently activate upstream steps in ATR signalling. Since these assays predominantly reflect the response of replicating phase cells, we first verified that CV1720 and control LBLs harbour a similar percentage of S phase cells (Figure S2). Pan-nuclear phosphorylation of H2AX (γH2AX) after replication fork stalling represents an ATR–specific damage response [24]. Strikingly, whilst exposure to 5 mM HU for 2 h resulted in an elevated number of cells staining positively for γH2AX in WT cells, this was not observed in either CV1720 or DK0064 (ATR–SS) cells (Figure 1B). We note that although previous studies have shown that ATM can be activated and phosphorylate γH2AX at DSBs arising following HU treatment in the absence of ATR due to enhanced fork collapse, this was not observed at 2 h post 5 mM HU exposure in these patient cells most likely due to residual ATR activity and/or the early times examined [26,27]. Chk1 represents an important ATR substrate required for G2/M checkpoint arrest. To examine Chk1 activation, we carried out Western Blotting using p-Chk1 antibodies. Following the same UV exposure conditions (2 h post 5J m\(^2\)) employed to examine G2/M checkpoint arrest, we
Figure 1. CV1720 cells show impaired ATR-dependent DNA damage responses. A) WT, DK0064 (ATR-SS), CV1720 (patient), CV1780 (patient’s mother) and CV1783 (patient’s father) cells were exposed to 5 J/m² UV and the mitotic index (MI) assessed 2 h post exposure. A greater than two fold decrease in mitotic index is observed in WT and both paternal cell lines but not in DK0064 (ATR-SS) or CV1720 (patient) cells. B) Cells were exposed to 5 mM HU for 2 h and the percentage of p-H2AX (∗-H2AX) positive cells assessed by immunofluorescence. Note that HU causes pan nuclear p-H2AX formation rather than defined foci as observed after exposure to ionising radiation. Thus, the percentage of ∗-H2AX positive cells was scored. C) Cells were exposed to UV (5 J/m²) and subjected to Western Blotting (WB) using p-Chk1 (p-Ser317) antibodies at 2 h. Chk1 expression was shown to be similar in WT and patient cells (lower panel). D) Cells were exposed to 3 mM HU for 2 h and whole cell extracts analysed by WB using FANCD2 antibodies. The ubiquitylation of FANCD2, detectable by a product with reduced mobility, is diminished in DK0064 (ATR-SS) and CV1720 cells compared to WT cells. E) Cells were exposed to 5 mM HU and examined for the percentage of cells showing ∗-53BP1 foci at 2 h post exposure. ∗-53BP1 foci formation is reduced in DK0064 (ATR-SS) and CV1720 cells compared to WT cells. F-I) The indicated cells were processed by WB using

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observed a pronounced p-Chk1 band in WT LBLs but not in CV1720 cells although Chk1 levels were similar in the two lines (Figure 1C). These results provide strong evidence that CV1720 show impaired ATR–dependent substrate phosphorylation.

A further ATR–dependent response is mono-ubiquitylation of FANCID2 following exposure to HU [20]. Mono-ubiquitinated FANCID2 can be detected by the presence of a slowly migrating isoform of FANCID2 generated post exposure to 3 mM HU. Whilst such a product was detected in WT cell extracts, it was absent in CV1720 and DK0064 (ATR–SS) cell extracts (Figure 1D). Finally, ATR also regulates the formation of 53BP1 foci following replication fork stalling via a Chk1-dependent process. We observed a failure to form 53BP1 foci following exposure to 5 mM HU in CV1720 and DK0064 (ATR–SS) LBLs in contrast to WT LBLs (Figure 1E), consistent with the diminished levels of p-Chk1 observed in CV1720 cells.

Collectively, these studies provide strong evidence that CV1720 cells are defective in an upstream step of the ATR–dependent signalling response defining them as distinct to the majority of previously characterised SS cell lines, which, though defective in UV-induced G2/M checkpoint arrest, are proficient in upstream steps of the ATR signalling response [25].

Reduced ATR and ATRIP protein expression in CV1720 cells

Given the overlapping cellular phenotype between CV1720 and DK0064 (ATR–SS) cells, we examined CV1720 cells for expression of ATR and ATRIP protein by Western Blotting. Strikingly, we observed markedly reduced levels of both ATR and ATRIP in CV1720 cells (Figure 1F). Since ATRIP stabilises ATR, this does not distinguish whether the primary defect lies in ATR or ATRIP and indeed a similar reduced level of ATR and ATRIP was observed in DK0064 (ATR–SS) cells (Figure 1G). Significantly, we observed reduced ATRIP and ATR in both parental LBLs (CV1780 and CV1783), which was approximately 50% of the level in WT LBLs (Figure 1G–1I).

Sequencing analysis reveals mutational changes in ATRIP in CV1720 cells

To examine whether the causal genetic defect in CV1720 lies in ATR or ATRIP, we carried out sequencing of the two genes. First, we undertook PCR-based gDNA sequencing of the 47 exons and neighbouring exon-intron boundaries of the human ATR gene from CV1720 cells and failed to observe any mutational changes likely to be of functional significance. Next, we undertook gDNA sequencing of ATRIP exons and observed a heterozygous mutational change, c.2278C>T, in exon12 which generated a stop codon predicting a truncated protein at position arginine 760 (p.Arg760*) (Figure S3). However, no mutational changes in any other exons were identified although we detected several novel intronic changes that could potentially impact on splicing (Table S1). Significantly, the c.2278C>T mutational change was observed as a heterozygous change in the patient’s mother but not in the father (Figure S3).

We also performed RT-PCR sequencing of ATRIP cDNA from CV1720 and both parents. These analyses revealed a low level of a smeared PCR product following amplification of the 5’ATRIP cDNA region using patient but not control cDNA (data not shown). Following multiple analyses, we found specifically that RT-PCR amplification using primers located in exons 1 and 4, reproducibly yielded a smeared product from CV1720 cDNA with discrete bands at 438 bp (the expected product size) and 325 bp whereas only the 458 bp product was observed using cDNA from WT cells (Figure 2A). Direct sequencing of the gel purified smaller (325 bp) and full-length (438 bp) RT-PCR products showed that the small fragment specifically lacked exon 2. Sequencing analysis of the RT-PCR product of CV1720 cDNA using the same primers revealed the predicted double sequence with the product lacking exon 2 being less than 50% of the product containing exon 2 (Figure 2B). It is notable that there were also some PCR products larger than the full length product although a discrete band was not evident. In sequencing the RT-PCR product, we observed some that harboured intron 2 sequences although these represented a minor product relative to that lacking exon 2. Collectively, these findings strongly suggested that there could be mis-splicing in CV1720 cells with loss of exon 2 being the major product.

To assess this further, qRT-PCR was undertaken using sets of primers that allow selective amplification of the WT and mutant products (c.2278C>T mutant as well as the mis-spliced product). The aim was to determine if the mis-spliced product originated from the paternal allele and if it impacted upon the transcript level. Primer pairs, P1 and P3C, located in exons 12 and 13, respectively, allow selective amplification of the WT (paternal) c.2278C allele whilst primers P2 and P3C selectively amplify the mutated (maternal) c.2278C>T allele (Figure 2C). As expected, the mutant (c.2278C>T)-allele-specific PCR product (right columns, red bars) was only detected in the patient and mother whereas the WT-specific PCR product (left columns, blue bars) was detected in all samples, demonstrating that the primers distinguished the two alleles (Figure 2C). The results also showed that the c.2278C>T and the WT (c.2278C) alleles were expressed at nearly equal levels (normalised against HPRT1) in the mother (compare blue and red bars labelled ‘mother’ in Figure 2C), suggesting that the c.2278C>T ATRIP mRNA is not subject to nonsense mediated RNA decay (NMD) (Figure 2C).

To evaluate the expression level of the mis-spliced ATRIP mRNA, we designed primers located at the exon 2/exon 3 boundary (primer P4) and within exon 3 (primer 6C) to allow selective amplification of the correctly spliced mRNA (Figure 2D); primers located at the exon 1/exon 3 boundary (primer P5) and within exon 3 (primer 6C) selectively amplify the mis-spliced mRNA. Whilst the correctly spliced product was amplified to similar (although slightly different) extents from father, mother and patient mRNA (compare the column heights, left panel in Figure 2D), the mis-spliced product was more abundant in the patient and father, suggesting that mis-splicing is a consequence of a mutational change linked to the paternal allele (compare the column heights, right panel in Figure 2D). Since we observed nearly equal expression levels of the wild type (c.2278C) and mutant (c.2278C>T) alleles in the mother (Figure 2C, compare the right and left panels), we considered that the PCR products derived from the mother using primers P4/P6C or P5/P6C would

ATRIP or ATR antibodies. MCM2 was used as a loading control. F shows the analysis of a range of protein levels for accurate comparison. CV1720 (patient) cells show markedly reduced ATR and ATRIP protein levels. G shows that both parental lines have approximately half the level of ATR and ATRIP compared to two WT cell lines. DK0064 (ATR–SS) and CV1720 cells, in contrast, have more dramatically reduced ATR and ATRIP protein levels. 50 ug protein was loaded. WT in all panels was GM2188. Patient, mother and father were as shown in panel A. H and I show the quantification of ATRIP and ATR protein levels from at least three independent WB experiments.

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be equally derived from the c.2278C and c.2278C>T alleles, which have, therefore, been depicted as equal sized contributions (shown in red or blue in mother columns in Figure 2D). Similarly, the mutant c.2278C>T allele is likely to be expressed at an equal level in the patient as in the mother (shown in red in patient columns in Figure 2D). Based on these assumptions, we estimated that the normally spliced WT mRNA is reduced to around 1/4 of the WT level in the patient and to 3/4 in the father (shown in blue in the left hand panel in Figure 2D). Assuming that the c.2278C>T allele is fully inactivated (see below), the patient

Figure 2. Identification of mutational changes in ATRIP in CV1720. A) Upper panel shows primer pairs used to distinguish cDNA products encompassing or lacking exon 2. Lower panel shows RT-PCR products obtained using the primers shown in the upper panel. RT-PCR from patient CV1720 generated a smeared product with a defined band of 458 bp, as observed in WT cells, and a weaker band of 325 bp. The latter band was not detected using cDNA from WT cells (MRC5). A similar single 458 bp band was obtained using the same primers with cDNA derived from a distinct wild type cell line (GM2188; data not shown). (B) Sequencing of the RT-PCR products derived from WT (MRC5) and patient (CV1720) cells. A double sequence pattern at the exon 2–3 boundary is observed using patient CV1720 cDNA. (C) Selective quantitative amplification of the WT or 2278C>T ATRIP alleles. Primers located in ATRIP exon 12 and 13 were designed to selectively amplify the WT (c.2278C) (P1 and P3C) versus the mutated (c.2278C>T) (P2 and P3C) alleles. The WT PCR product is shown in blue and the c.2278C>T PCR product in red. The exon 12 mutated allele is only observed in the patient and mother cDNA whilst the WT allele is observed in the patient, mother and father cDNA although the level is reduced in the patient and mother. (D) qRT-PCR analysis of ATRIP splicing variants from patient CV1720 and parental cells. qRT-PCR analysis of the level of the normally spliced (encompassing exons 1-2-3) and the aberrantly spliced (∆exon2) ATRIP cDNA in the patient and parent cells. PCR primers were designed at the exon2-exon3 or exon1-exon3 boundaries to selectively amplify the splicing variants. Transcripts from HPRT1 were used as a quantification control. The correctly spliced transcript from the paternal allele of the patient (wild type c.2278C, blue fraction in the cumulative bar labelled, 'patient', at the left panel) was estimated to be ~25% of the normal level. (E) The mis-spliced paternal allele is subject to nonsense mediated mRNA decay (NMD). Cycleave-qPCR confirmed that the ATRIP c.2278C>T mutant allele was expressed exclusively in the patient and the mother. The ATRIP exon12-13 fragment was amplified with PCR primers P7/P8 as shown in the figure. A set of fluorescent probes were used to distinguish the WT versus c.2278C>T allele (probe1 and probe2, respectively). In the patient, the paternal mRNA transcript level (emerald lines) is low because of NMD (top left). Puromycin treatment eliminated the NMD and the paternal transcript level returned to the normal level. In all panels WT represented MRC5, patient was CV1720 and parents were as shown in Figure 1A.

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therefore has around 25% of ATRIP activity compared to a normal individual.

The findings above suggested that the mis-spliced mRNA, which generates an out of frame cDNA, is subject to NMD. To examine this and substantiate our findings, qRT-PCR was also carried out using fluorescent cycleave probes with or without exposure to puromycin, an antibiotic which prevents NMD (Figure 2E) [29]. Primers (P7 and P9) and fluorescent probes (probe 1 and 2) were designed to allow amplification of a product encompassing exon 12–13 that distinguished the maternal (probe 2) from the paternal (probe 1) allele. We confirmed detection of the c.2278C >T allele exclusively in the patient and mother as well as the WT allele in all samples (Figure 2E). In the mother, the wild type (c.2278C) and mutant (c.2278C>T) signals were detected at equal levels regardless of whether puromycin was added, indicating that the alleles are equally expressed and are not subjected to NMD. In patient CV1720, the WT product was reduced relative to the mutant product in the absence of puromycin but was at similar levels in the presence of puromycin (Figure 2E). These findings are consistent with the notion that the mRNA expressed from the parental allele is aberrantly spliced and partially subject to NMD. Perhaps surprisingly, we did not detect any obvious difference of the WT product following puromycin treatment in the father; however, in this case, we anticipate a 25% decreased product, which is unlikely to be detected without an internal control. However, despite this, there was evidence for abnormal splicing in the paternal cDNA from analysis of the PCR products spanning exons 1–3 (Figure 2B, 2D).

Finally, to gain insight into the basis underlying mis-splicing, we sequenced introns 1 and 2 from the patient, mother, and father and identified a previously unreported mutational change in intron 2, 13 bp from the intron-exon 2 boundary in the patient and paternal gDNA (Table S1). However, given the modest impact on splicing we did not attempt to examine whether this represented the causal mutational change affecting splicing.

Arg760* ATRIP does not promote ATR–dependent G2/M arrest and reduces ATR–ATRIP interaction

It is likely that ATRIP c.2278C>T causes an impacting mutational change since the low levels of ATRIP protein (10–20% WT levels) in CV1720 cells suggest that p.R760* ATRIP is unstable (given that the mRNA level of this allele is normal). To substantiate that p.R760* expression impairs the ATR–dependent response to DNA damage, we examined whether its expression could complement the G2/M checkpoint defect of CV1720 cells. We also examined whether p.R760* might exert a dominant negative impact (since this represented a possible explanation for the low ATRIP protein level in CV1720 cells). The c.2278C>T mutational change was introduced into ATRIP cDNA by site directed mutagenesis. CDNA encoding WT ATRIP and/or R760* ATRIP was transiently transfected into LBLs and G2/M checkpoint arrest examined at 2 h post exposure to 5 Jm-2 UV. Consistent with previous findings, WT but not CV1720 cells showed a G2/M checkpoint arrest (Figure 3A). Whilst transfection with WT ATRIP cDNA completely rescued the G2/M checkpoint defect of CV1720 cells, no correction was observed in CV1720 cells following expression of c.2278C>T ATRIP cDNA (encoding R760* ATRIP). Surprisingly, expression of WT ATRIP cDNA also corrected the G2/M checkpoint defect in DK0064 (ATR–SS) cells, which we propose could result from elevated ATRIP expression causing stabilisation of residual ATR protein, since ATR–SS cells have low ATR and ATRIP expression. Significantly, c.2278C>T ATRIP cDNA was unable to rescue ATR–SS cells.
Checkpoint arrest after exposure to ionising radiation was activated normally. Additionally, we examined the phosphorylation of a range of ATR substrates following exposure to 0.5 mM HU and observed impaired phosphorylation in both 27-4BI and 19-8BI cells (Figure 5B). Collectively, these functional data substantiate a deficiency in the ATR–dependent DNA damage response in LBLs from these two cases. Thus, we conclude that both patients represent further ATR–SS patients.

Discussion

Although the first causal defect for SS was identified as ATR in 2003, further patients with mutations in ATR have not been reported [7]. SS patients are characterised by microcephaly and growth delay, features also observed in other microcephalic, primordial dwarfism syndromes including MOPD type II and MGS. Given that all ATR–SS patients to date share consanguinity, there are limitations in defining the spectrum of clinical features conferred by ATR deficiency to support a clinical distinction between ATR–SS and related disorders such as MOPD type II and MGS as well as other sub-classes of SS [1–3,31].

Here, we describe the novel identification of a patient mutated in ATRIP, the binding partner of ATR. Thus, we identify ATRIP as a new causal gene for SS. The mutational change in one ATRIP allele lies within a region previously suggested to be required for interaction with ATR, which is consolidated by our work [21,30]. We demonstrate that the second allele is abnormally spliced causing a reduction in ATRIP mRNA from that allele. qRT-PCR analysis suggested that there could be 25% residual WT ATRIP expressed in the patient cells. Consistent with this, we routinely observed 10–20% of WT ATRIP protein in CV1720 cells by Western Blotting, although the level was variable between preparations. Although not examined in detail, there appeared to be a correlation between proliferation status and ATRIP levels, with the levels decreasing as proliferation slowed. Thus, differences in the proliferative state of cells at the time of analysis may underlie the apparent difference between Western Blotting and qRT-PCR analysis. Notwithstanding some limitations in quantification, the patient clinical features were marked despite ~10–20% residual WT ATRIP expression. Similarly, in patient DK0064, residual ATR protein can be readily detected [7]. Thus, we conclude that reduced but detectable levels of ATR/ATRIP protein can confer a clinical phenotype.

Additionally, we identify two further SS patients with ATR mutations in two unrelated families native to the UK. Interestingly, despite being unrelated, 27-4BI and 19-8BI carry the same compound heterozygous mutations, possibly representing founder mutations in the UK population.
All four ATR/ATRIP patients displayed severe microcephaly and growth delay (Table 1). All patients also displayed micrognathia, receding forehead, dental crowding and microtia with small or absent lobes (Figure 4A). Interestingly, an MRI scan of the ATRIP–SS patient revealed an abnormally small pituitary with absent fossa, which could contribute to the delayed growth observed (Figure S1). In distinction to the original ATR–SS patient (DK0064), patients 27-4BI and 19-8BI showed more marked skeletal abnormalities including digital features and aberrant patellae suggesting that ATR deficiency can have a detrimental impact on bone development (Table 1, Figure S4) [32]. Interestingly, aberrant patellae is a clinical feature commonly exhibited by MGS patients suggestive of a biological overlap between the ATR checkpoint pathway and the replication machinery during skeletal development and maintenance. In keeping with this, characterisation of a mouse model harbouring the same mutational change identified in the original ATR–SS patient (DK0064) revealed marked bone abnormalities including kyphosis and osteoporosis [32]. Our findings suggest that ATR–ATRIP SS shows more overlap with MGS than previously recognised (Table 2). However, whereas ATR–ATRIP SS patients tend to have very marked microcephaly, growth delay, dental crowding, small ears and less severe skeletal abnormalities, the spectrum for MGS tends to be less marked microcephaly and growth delay but a striking impact on skeletal development. Nonetheless, there does not appear to be an absolute clinical divide between these two disorders. Significantly, these overlapping clinical features could reflect the fact that both ATR/ATRIP

Figure 4. Patients 27-4BI and 19-8BI have reduced ATR and ATRIP expression and mutations in ATR. A) Photographs of patient included with informed consent of parent. B) Cell extracts (50 µg) from LBLs derived from WT (IM257), patient 27-4BI or patient 19-8BI were immunoblotted using the indicated antibodies. Reduced expression of ATR was observed in both patients. 27-4BI also had reduced ATRIP expression. C) Structure of ATR showing the site of the mutations identified and the UME domain. D) The UME domain is conserved between species and the methionine residue within this domain is conserved in yeast.
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and the origin licensing complex play an essential role in promoting efficient replication and recovery from fork stalling, which may be vital during developmental stages involving rapid replication [13].

In summary, we provide the first report of a SS patient with mutations in ATRIP, defining a further novel genetic defect for this disorder, and describe two additional patients native to the UK, with mutations in ATR. The description of multiple ATR–ATRIP patients allows us to define a spectrum of clinical features conferred by ATR–ATRIP mutations. The clinical characteristics include severe microcephaly and growth delay, small or absent ear lobes, micrognathia and dental crowding. In addition, the novel ATR–mutated cases described here expand the clinical impact of impaired ATR–function to include more marked skeletal involvement.

Figure 5. LBLs from patient 27-4BI and 19-8BI showed impaired ATR–dependent damage responses. A) 27-4BI cells were examined for their ability to activate G2/M checkpoint arrest at 4 h following exposure to 7 J/m² UV. In contrast to WT cells (GM2188), no significant arrest was observed in 27-4BI cells. The checkpoint response to ionizing radiation, which is ATM rather than ATR dependent, was normal. B) LBLs derived from patients 27-4BI and 19-8BI were examined for their ability to phosphorylate the indicated ATR substrates at 1 h following exposure to 0.5 mM HU. WT represents IM257. 27-4BI and control LBLs have a similar cell cycle profile demonstrating that the lack of ATR substrate phosphorylation cannot be attributed to the lack of S phase cells (Figure S2).

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Methods

Ethics Statement

Ethical approval for the research was granted by the School of Life Sciences Research Governance Committee, University of Sussex. Informed consent was obtained and clinical investigations were conducted according to the principles expressed in the Declaration of Helsinki. Patient material was gathered under conditions of the Human Tissue Authority (HTA licence number 12119).

Patients and cell lines

CV1720 is a SS patient of Gujarati-Indian origin. Patients 27-4BI and 19-8BI are English. The clinical features are described in
Table 1. Lymphoblastoid cell lines (LBLs) were derived from blood following EBV transformation. WT LBLs were GM2188 or LB197 as indicated. All LBLs were grown in RPMI medium supplemented with 10% foetal calf serum, penicillin, and streptomycin. Transfection with ATRIP cDNA was with Genejuice Transfection Reagent (Novagen, Merck Millipore, UK) following a nitrocellulose membrane for protein detection.

<table>
<thead>
<tr>
<th>Number of patients</th>
<th>ORC1 - MGS</th>
<th>Pre-RC MGS</th>
<th>ATR/ATRIP SS</th>
</tr>
</thead>
<tbody>
<tr>
<td>OFC (cm)*</td>
<td>–5.4 to –11 SD</td>
<td>+1.7 to –5.0 SD</td>
<td>–10 to –12 SD</td>
</tr>
<tr>
<td>Height (cm)*</td>
<td>–4.5 to –9.6 SD</td>
<td>–0.4 to –6.4 SD</td>
<td>–5 to –8 SD</td>
</tr>
<tr>
<td>Weight (kg)*</td>
<td>0.8 to –11 SD</td>
<td>–0.3 to –9.9 SD</td>
<td>–3.3 to –8 SD</td>
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<tr>
<td>Intellectual disability</td>
<td>Ranges from none to mild/moderate</td>
<td>None</td>
<td>Developmental delay (2/4)</td>
</tr>
<tr>
<td>MRI</td>
<td>Normal in 2 patients examined</td>
<td>NA</td>
<td>Generalised cerebral atrophy, delayed myelination, abnormal gyration (2 patients examined)</td>
</tr>
<tr>
<td>Other</td>
<td>High pitched voice (1/10), full lips (7/10), cryptochordism (2/4 examined), mammmary hypoplasia (2/2 examined), feeding and respiratory problems during infancy (8/10)</td>
<td>Full lips (14/25), cryptochordism (7/14 examined), mammmary hypoplasia (8/8 examined), feeding (20/25) and respiratory (9/25) problems during infancy</td>
<td>Dental crowding (4/4), feeding and respiratory problems during infancy (1/4)</td>
</tr>
</tbody>
</table>

*standard deviations from the age-related normal population mean, NA = not assessed.

MGS data from [13,34] [33,34]. doi:10.1371/journal.pgen.1002945.t002

**Table 2.** MGS and Seckel syndrome patient phenotypes.

**ORC1 - MGS** | **Pre-RC MGS** | **ATR/ATRIP SS**
---|---|---
Number of patients | 10 | 25 | 4
OFC (cm)* | –5.4 to –11 SD | +1.7 to –5.0 SD | –10 to –12 SD
Height (cm)* | –4.5 to –9.6 SD | –0.4 to –6.4 SD | –5 to –8 SD
Weight (kg)* | 0.8 to –11 SD | –0.3 to –9.9 SD | –3.3 to –8 SD
Intelligence disability | Ranges from none to mild/moderate | None | Developmental delay (2/4)
MRI | Normal in 2 patients examined | NA | Generalised cerebral atrophy, delayed myelination, abnormal gyration (2 patients examined)
Other | High pitched voice (1/10), full lips (7/10), cryptochordism (2/4 examined), mammmary hypoplasia (2/2 examined), feeding and respiratory problems during infancy (8/10) | Full lips (14/25), cryptochordism (7/14 examined), mammmary hypoplasia (8/8 examined), feeding (20/25) and respiratory (9/25) problems during infancy | Dental crowding (4/4), feeding and respiratory problems during infancy (1/4)

**Immunofluorescence for analysis of γH2AX and 53BP1 staining**

Cells were cytopsorn onto slides, fixed with 3% formaldehyde for 10 min and permeabilized in 0.5% Triton-X100. After antibody treatment and staining with 4,6-diamidino-2-phenylindole (DAPI), coverslips were mounted in Vectashield mounting medium (Vector Laboratories, Burlingame). Samples were incubated with primary antibodies for γH2AX (Millipore, Billerica) or 53BP1 (Bethyl, Montgomery). Secondary antibodies were from Sigma (St. Louis).

**Western blotting**

Cells were lysed for one hour in IPLB (50 mM Tris-HCl, 150 mM NaCl, 2 mM EDTA, 2 mM EGTA, 25 mM NaF, 25 mM β-glycerolphosphate, 0.1 mM NaOrthovanadate, 0.2% Triton X-100, 0.3% NP-40, plus protease inhibitor cocktail (Roche, Basel) at 4°C, centrifuged at 15,000 rpm for 10 minutes. The soluble fraction was subjected to SDS-PAGE and transferred to a nitrocellulose membrane for protein detection.

**Antibodies against ATR, CHK1 (FLA76) and MCM2 (N19) were from Santa Cruz (Santa Cruz). Anti-FANC2D, ATRIP and phospho-Chk1 (Ser317) antibodies were from Novus (Littleton), Bethyl (Montgomery), and Cell Signaling (Beverly, Woburn), respectively.**

**G2/M checkpoint arrest**

Cells were exposed to 5 or 7 Jm⁻² UV, or 3Gy ionising radiation and incubated for 2 or 4 hours (as indicated) in complete medium containing 0.2 ug/ml Colcemid (Invitrogen, Carlsbad), followed by processing for immunofluorescence as detailed above. Mitotic cells were detected by a-Histone H3-pSer10 antibodies (Millipore, Billerica) and cells were counterstained with DAPI.
Supporting Information

Figure S1 Photograph of limbs and MRI scan of patient CV1720. Left hand photograph showing hands and feet. Right hand photograph shows an MRI scan where a small pituitary is evident. (TIF)

Figure S2 Cell cycle analysis of WT, DK0064 (ATR–SS), 27-4BI, and CV1720 patient LBLs. A) Asynchronous growing cultures of WT, DK0064 (ATR–SS), 27-4BI and CV1720 patient cells were pelleted, fixed in 70% ice-cold ethanol and stained with propidium iodide prior to FACS analysis. Populations were gated and the proportion of cells in G1, S and G2/M phases of the cell cycle measured. B) WT, DK0064 (ATR–SS), 27-4BI and CV1720 patient cells were treated with nocodazole for 16 h and then fixed and analysed as in a). C) Asynchronous growing cultures of WT, DK0064 (ATR–SS), 27-4BI and CV1720 patient cells were pulse-labelled with 50 μM BrdU for 1 h. Cells were then fixed in 70% ice-cold ethanol and processed for BrdU FACS analysis as described in Bicknell et al, 2011 [13]. The proportion of cells in S phase were gated and measured. Each graph represents the mean of three independent experiments. The error bars represent the standard deviation. (TIF)

Figure S3 Identification of a truncating mutation in ATRIP in patient, CV1720. Genomic DNA sequencing of ATRIP exons showed that patient CV1720 and the unaffected mother, CV1780, are heterozygous for a c.2278C>T mutational change in exon12 of ATRIP. The father has a WT sequence at this site. 2278C>T generates a primary stop codon predicting a truncated protein at position arginine 760 (p.R760*). WT sequence shown in blue, the mutation is shown in Red. (TIF)

Figure S4 Photographs of patients 27-4BI and 19-8BI. A) Shows abnormal digits of patient 27-4BI. B) Copper beaten appearance of skull of patient 19-8BI. C) Frontal and Lateral view of left knee of patient 19-8BI showing an absence of ossification of the patella. (TIF)

Figure S5 Mutational changes observed in ATRIP in patients 27-4BI and 19-8BI. A) c.3477G>T mutational change in patients 27-4BI and 19-8BI. RT-PCR sequencing revealed a heterozygous 3477G>T mutational change in both patients causing an amino acid substitution, p.Met1159Ile, which lies within a conserved UME (NUC010) domain of ATR. B) A double sequence was observed at the boundary between exons 40 and 41 in both patients. Sequencing showed that the double sequence was caused by insertion of a 142 bp region from intron 40. C) A C to G mutational change was observed in intron 41 of both patients converting the sequence CAGCT to CAGGT, a splice site. The insertion causes a frameshift and a stop codon at p.Val2300-Gly8*75. D) Diagram showing the likely origin of the insertion observed at the exon 40/41 boundary. Sequencing of intron 40 revealed a C>G mutation as indicated creating a cryptic splice site causing splicing of exon 40 to the indicated intrinsic sequence (which represents an Alu repeat sequence). Thus one ATR allele of the patients harbours a 142 nucleotide insertion between exons 40 and 41. Exons 40 and 41 are highlighted in green and the inserted intrinsic sequence is shown in red. The intronic C>G change is highlighted in red. The insertion causes a frameshift and a stop codon at c.6976 in exon 41. (TIF)

Table S1 The table shows the position of single nucleotide polymorphisms identified in intron 1 and 2 in the patient and parental genomic DNA. * The contig position is defined as the position of the single nucleotide variant (SNV) on the contig (NT_002517.17) when counting from the first base (base position = 1). **rs# is the NCBI’s reference SNP ID. *** minor allele (indicated as a base) and its frequency (MAF) (second most frequent allele) in a default global population reported in dbSNP database (1000 Genome phase 1, May 2011). N.A. not available. (DOCX)

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Author Contributions

Conceived and designed the experiments: TO GSS AMRT PO AJ. Performed the experiments: SW TS SL GC MM NM YN. Analyzed the data: TO GSS AMRT MO PAJ. Contributed reagents/materials/analysis tools: EH KP MS PJ PB MB. Wrote the paper: PAJ. Aided in writing the paper: MO GSS TO AMRT.

References