Identification of a functionally significant tri-allelic genotype in the Tyrosinase gene (TYR) causing hypomorphic oculocutaneous albinism (OCA1B)

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Oculocutaneous albinism (OCA) and ocular albinism (OA) are inherited disorders of melanin biosynthesis, resulting in loss of pigment and severe visual deficits. OCA encompasses a range of subtypes with overlapping, often hypomorphic phenotypes. OCA1 is the most common cause of albinism in European populations and is inherited through autosomal recessive mutations in the Tyrosinase (TYR) gene. However, there is a high level of reported missing heritability, where only a single heterozygous mutation is found in TYR. This is also the case for other OCA subtypes including OCA2 caused by mutations in the OCA2 gene. Here we have interrogated the genetic cause of albinism in a well phenotyped, hypomorphic albinism population by sequencing a broad gene panel and performing segregation studies on phenotyped family members. Of eighteen probands we can confidently diagnose three with OA and OCA2, and one with a PAX6 mutation. Of six probands with only a single heterozygous mutation in TYR, all were found to have the two common variants S192Y and R402Q. Our results suggest that a combination of R402Q and S192Y with a deleterious mutation in a ‘tri-allelic genotype’ can account for missing heritability in some hypomorphic OCA1 albinism phenotypes.

Oculocutaneous albinism (OCA) and X-linked ocular albinism (OA) are inherited disorders of melanin biosynthesis which result in varied levels of hypopigmentation of skin, hair, and ocular tissues1. Characteristic ophthalmic features include reduced visual acuity, nystagmus, strabismus, and photophobia. Closer examination may reveal foveal hypoplasia (abnormal retinal development), asymmetry of visual evoked potential (VEP) responses, and iris transillumination1. Foveal hypoplasia for instance, can be determined using Spectral-Domain Optical Coherence Tomography (SD-OCT) and then graded on a scale of 1–4 (Thomas et al.2), and the asymmetry of visual-evoked potentials documents the excessive decussation at the optic chiasm seen in albinism1. Partial phenotypes are described widely in the literature in which some features are present but others are lacking (e.g. nystagmus or foveal hypoplasia), however, phenotyping methods have varied significantly and the partial phenotype has never before been described in detail4–6. Current management of albinism focusses on correction of any

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phenotype. OCA1 has a mixed phenotype and is further split into OCA1A and OCA1B. OCA1A describes a 40% reduction of tyrosinase enzymatic activity\(^{24}\). Multiple OCA1 studies have shown the R402Q allele is strongly associated with albinism patients with only one mutation\(^{12, 17, 20}\). It is also important to note that a variety of techniques have been employed to screen for tyrosinase gene mutations in these studies and no method has 100% sensitivity. Six genes involved in melanin biosynthesis pathway are known to cause forms of OCA and OA: TYR (tyrosinase), OCA2, TYRP1 (tyrosinase-like protein 1), SLC45A2 (solute carrier family 45 member 2), SLC24A5 (solute carrier family 24 member 5), and C10orf11 (chromosome 10 open reading frame 11) accounting for OCA subtypes 1–4 and 6–7 respectively, and GPR143 accounting for OA\(^1\), see Table 1. All of the OCA subtypes are understood to be inherited as autosomal recessive disorders but the subtypes are heterogeneous in pigmentation phenotype\(^6\). OCA1 has a mixed phenotype and is further split into OCA1A and OCA1B. OCA1A describes complete loss of tyrosinase activity (previously described as ‘tyrosine negative’ albinism) and is characterised by an apparent total lack of pigment. Some tyrosinase function is retained in OCA1B, allowing pigment to accumulate and generate a phenotype of minimal to near normal skin pigmentation, as is also the case for the other described OCA and OA phenotypes\(^7, 8\). Phenotypes of partial OCA also overlap with those seen in patients with dominant mutations in the PAX6 gene, which is involved in ocular development, where a variety of phenotypes have been described including lveal hypoplasia, iris trans-illumination and nystagmus\(^7\).

As the most severe form of OCA, OCA1A is often recognised in early infancy. King et al. proposed that white hair from birth can be used to predict OCA1\(^{15}\), with 85% of patients identified in this way testing positive for pathogenic TYR mutations. However, 15% of OCA cases identified in this way had no accountable genetic mutation, and 29% of those confirmed as OCA1 had only one identifiable TYR mutation\(^6\). It is widely recognised that the OCA genes do not account for all non-syndromic cases, as many as 30% of OCA1A occurrences have an unknown genetic origin\(^{10, 11}\) and this percentage may be higher for cases of partial albinism\(^{12}\). It is also important to note that a variety of techniques have been employed to screen for tyrosinase gene mutations in these studies and no method has 100% sensitivity. An individual’s pigmented phenotype depends on polymorphisms in many genes, including polymorphisms in the OCA genes\(^3, 4\). Ethnic background may play a large role in an individual’s susceptibility to the albinism phenotype, with hypomorphic mutations having a more damaging effect on a less active pigmentation pathway\(^4\). It has been suggested that inheritance of OCA2 is not purely recessive, with the example of haploinsufficiency noticeably affecting skin complexion in a Hispanic family, arguably due to the already fair skin tone\(^5\). It has also been suggested that a synergistic interaction between genes throughout the pigment pathway may exist in albinism phenotypes, evidenced by one family exhibiting an OCA2 phenotype that is modified by a mutation in the gene for OCA3\(^4\) and a correlation between OCA2 and MC1R variants in a small albinism cohort\(^8\). The quantitative effect of pigmentation also has relevance to OCA1B, particularly the notion of autosomal recessive ocular albinism (AROA), an arbitrary characterisation that has been used previously to describe cases with clinically mild OCA1B\(^9, 10\).

AROA sparked a debate over the possible pathogenicity of two TYR polymorphisms, rs1126809 (p.R402Q) and rs1042602 (p.S192Y), common in Caucasian populations with allele frequencies ~28–36%\(^{15}\). Functional studies have shown the R402Q polymorphism produces a thermolabile enzyme, retained by the cells endoplasmic reticulum, with a 75% reduction in catalytic activity compared to the wild-type\(^{15, 22, 23}\); and S192Y results in a 40% reduction of tyrosinase enzymatic activity\(^{24}\). Multiple OCA1 studies have shown the R402Q allele is strongly associated with albinism patients with only one mutation\(^1, 2, 7, 20\).

R402Q has been proposed as a causal variant, though only when inherited on the trans allele to a null activity TYR mutation\(^{15, 20}\). However this was disputed with evidence of no OCA phenotype in the parents of affected probands even when they carried a combination of null mutation and R402Q\(^{25}\). This has led to the question of whether it is possible for an additional variant to be necessary for manifestation of the ocular phenotype. The combination of two common variants may produce a reduction in TYR activity that, when co-inherited with a deleterious TYR mutation, provides sufficient loss of activity to cause an albino phenotype\(^{15, 16}\). A similar tri-allelic hypothesis has been demonstrated in Bardet-Biedl syndrome\(^{26}\), but is yet to be demonstrated in albinism.

In this study, we have sequenced all the known albinism genes in patients with possible hypomorphic albinism phenotypes, identified through detailed ocular phenotyping in a tertiary eye clinic. Proband with some, but not all of the typical cutaneous and ocular features of OCA1A were defined as having a likely hypomorph albinism phenotype. For the first time, we investigate common variants in tri-allelic pattern of inheritance using detailed phenotyping and segregation studies in relatives to identify the causative genotype.

| HGNC symbol | HGNC name | Albinism subtype | Mode of inheritance |
|-------------|-----------|-----------------|--------------------|
| TYR         | Tyrosinase | OCA1A           | Autosomal recessive |
| OCA2 (P gene) | OCA2 melanosomal transmembrane protein | OCA2 | Autosomal recessive |
| TYRP1       | Tyrosinase related protein 1 | OCA3 | Autosomal recessive |
| SLC45A2     | Solute carrier family 45 member 2 | OCA4 | Autosomal recessive |
| ---          | Chromosomal location 4q24 | OCA5 | Autosomal recessive |
| SLC24A5     | Solute carrier family 24 member 5 | OCA6 | Autosomal recessive |
| C10orf11    | Chromosome 10 open reading frame 11 | OCA7 | Autosomal recessive |
| GPR143      | G protein-coupled receptor 143 | OA1 | X-linked recessive |

Table 1. Table to describe HGNC approved gene names associated with the subtypes of OCA and OA. OCA5 has been attributed to a chromosomal location but does not yet have an associated gene\(^46\).
Methods

Patients were recruited following the tenets of the declaration of Helsinki, informed consent was obtained and the research was approved by the Southampton & South West Hampshire Research Ethics Committee.

We investigated the genetic cause of eighteen probands categorized as having hypomorphic albinism. Probands were identified from a regional paediatric nystagmus clinic. All patients seen in this clinic underwent detailed phenotyping of skin and hair tone in context of family pigmentation, orthoptic examination, anterior and posterior segment examinations on a slit-lamp biomicroscope, electrodiagnostics including an electroretinogram (ERG) and visual evoked potential (VEP), and optical coherence tomography (OCT) of the macular using either a Leica OCT system or a Spectralis OCT (Heidelberg Engineering). Eye movement recordings were made on an EYELink1000 + (SR research) eye tracker and refraction was measured. Saliva was collected and DNA extracted using Oragene-DNA kit (OG-575) (DNA Genotek).

Probands with at least two phenotypic features of albinism (skin and hair pigmentation deemed to be low within the family context/nystagmus/foveal hypoplasia/VEP crossing/iris transillumination) as determined by a consultant ophthalmologist (JES), were chosen from a larger database containing approximately 300 probands with albino and/or nystagmus phenotypes. Probands were additionally excluded if they had complete characteristics of OCA1A or where DNA quality was poor.

The DNA samples were enriched using the TruSight One capture platform (Illumina 5200 Illumina Way San Diego, California USA). TruSight One has been dubbed a “clinical exome”, covering 4813 genes associated with disease-causing mutations. The panel targets and captures most of the coding regions of OCA genes 1–4 & 6, the OA1 gene, all syndromic albinism genes and PAX6, coverage of genes is shown in Supplementary Table 1. Prepared libraries underwent paired-end sequencing on an Illumina NextSeq 500 machine.

Next generation sequencing (NGS) data was aligned against the human reference genome (hg19) using Novosalign (v2.08.02). The mean read depth across all samples was 167 (Supplementary Table 1) with 97.2% of all target regions achieving a depth of 20X or greater. Variant calling was performed using SAMTools v0.1.19 and variant annotation using ANNOVAR against RefSeq transcripts. Additional annotation was applied using the Human Gene Mutation Database, HGMD.

Prepared libraries underwent paired-end sequencing on an Illumina NextSeq 500 machine. The hypomorphic albinism phenotype varied in both ocular phenotype and pigment level between probands and between family members. For example proband and mother in family 3 both have a phenotype consistent with partial albinism, however the proband exhibits a severe loss of cutaneous pigment but no iris transillumination, whereas the cutaneous pigment in the proband’s mother is within that of the family context but ocular investigations revealed trans-illumination defects. The level of foveal hypoplasia also varied between patients and within families. Example OCT images taken from the cohort are in Fig. 1, demonstrating the broad range of foveal developmental anomalies identified.

NGS data for OCA genes 1–4 & 6, the OA1 gene, and PAX6 were initially filtered using predictive scores from SIFT and PolyPhen. In silico pathogenicity prediction tools SIFT (<0.05), PolyPhen2 HumVar (possibly damaging and probably damaging) and GERP++ (≥2). SIFT predicts pathogenicity of missense mutations based on homology, PolyPhen2 HumVar predicts pathogenicity based on conservation and protein structure/function and GERP++ measures evolutionary constraint. The six probands with only a single heterozygous mutation were further investigated. Sanger sequencing was used to confirm and segregate each TYR variant in probands and family members, primers used are listed in Supplementary Table 3. Primers designed by Chaki et al. were used to for amplification of TYR exon 4 to avoid amplification of the highly homologous TYRI gene.

Multiple ligation-dependent probe amplification (MLPA) was carried out for the TYR and OCA2 genes as according to the manufacturer’s instructions with the current SALSA MLPA P325 OCA2 probe mix at the time of testing (MRC-Holland, the Netherlands). Partial albinism probands and control individuals were compared. Subsequent data were analysed using the MLPA analysis software of the GeneMarker (version 1.85) software (SoftGenetics, USA).

Results

Diagnosis of hypomorphic albinism. The hypomorphic albinism phenotype varied in both ocular phenotype and pigment level between probands and between family members. For example proband and mother in family 3 both have a phenotype consistent with partial albinism, however the proband exhibits a severe loss of cutaneous pigment but no iris transillumination, whereas the cutaneous pigment in the proband’s mother is within that of the family context but ocular investigations revealed trans-illumination defects. The level of foveal hypoplasia also varied between patients and within families. Example OCT images taken from the cohort are in Fig. 1, demonstrating the broad range of foveal developmental anomalies identified.

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Segregation of the OCA1 tri-allelic genotype. We further investigated the single TYR variants in both probands and family members (families 4–7, 12 and 18 in Table 3) using Sanger sequencing to confirm and determine segregation of variants. In total, twenty probands and family members were phenotyped and genotyped, results in Table 3. The phenotyping results of these six families suggests a total nine cases of partial albinism (six probands and three affected family members). Sanger sequencing confirmed the predicted causal variants in probands and revealed variants segregated with affected family members in every case, with three unaffected family members as carriers.

To explore the apparent missing heritability in these cases we investigated the potential pathogenicity of common variants R402Q and S192Y. The NGS data was examined in probands with TYR mutations. All six probands were found to have both common variants. These variants were confirmed in probands with Sanger sequencing and variant segregation was determined across available members of the six pedigrees, shown in Fig. 2. The combined presence of both common polymorphisms and a putative TYR mutation in a tri-allelic genotype segregates with affected family members.

It can be deduced that the R402Q variant is on the trans allele to the deleterious TYR mutation in probands 4, 5, 7 and 12. In family 4 we can also be certain the S192Y variant is on the trans allele. The mother of proband 18 has both nystagmus and foveal hypoplasia, yet does not have the same deleterious TYR mutation as her son.

Discussion
We have combined high resolution phenotyping, a broad NGS technique, segregation analysis and MLPA studies in a cohort of presumed partial albinism patients. This allows us the opportunity to perform a detailed genotype-phenotype correlation in this group of patients for the first time. In this study we identified one novel variant in the PAX6 gene, a novel frameshift variant in the GPR143 gene, two novel variants in the OCA2 gene (both in probands 3), five previously reported variants in OCA2, three novel and four previously reported variants in the TYR gene, and one previously reported variant in TYRP1 in eighteen probands. When combined, these variants provide a convincing genetic diagnosis for only 22% of our original hypomorphic albinism cohort if those with missing variants, in a presumed recessive condition (OCA1), are excluded.

The novel variant in GPR143, c.485delG, causes a frameshift mutation likely resulting in ocular albinism in proband 10. Of the six different mutations found in OCA2; N465D, V419I, Y342C and L650V have been reported previously in association with albinism. The variants R536C and W274C are both predicted to be deleterious by SIFT, PolyPhen2 and GERP++, described in Table 2.

The probands revealed seven different mutations in the TYR gene: V177F, c.1467dup, c.505_507del, C244Ter, R422W, R402Ter and P406L. The mutation V177F has been previously reported in an albinism cohort. TYR
NM_001300984, OCA2 prediction score was not available at the time of analysis. Gene accessions number: PAX6, C10orf11.

R422W has been reported as disease causing, however functional studies of this mutation have conflicting results. Mondal et al. assayed the tyrosine hydroxylase and DOPA oxidase activity for a partial OCA1 phenotype. codon is considered highly deleterious. It is likely that further functional analyses are necessary to produce a curated list of mutations for accurate genetic diagnosis.

Six probands within our cohort were found to have single TYR variant previously identified in albinism patients, but no variant in another known gene. As there is no functional evidence for the variants in family 5 and family 6 there remains the possibility of another causal gene mutation. It has been suggested that this high level of missing heritability could be due to mutations in the TYR promoter or an interacting distal gene enhancer. Notably, all six had also inherited R402Q and S192Y common TYR variants producing a tri-allelic genotype. It is likely that further functional analyses are necessary to produce a curated list of mutations for accurate genetic diagnosis.

The common variant R402Q is located in exon 4, near to the CuB catalytic site, and produces a thermolabile enzyme, but it has been argued that the reduction of tyrosinase activity is not enough to produce a phenotype. The controversy over the R402Q variant stems from a paper by Oetting et al. which argues that segregation of R402Q with a known pathogenic variant on the homologous allele does not confer albinism. The variant S192Y is located in the CuA catalytic site of tyrosinase and has been shown to lower enzymatic activity independently to R402Q. Previous studies have had stringent criteria for an OCA1 phenotype (white hair and skin and translucent irides from birth), whereas, here we have considered hypomorphic presentations that do not appear as severe but result in ocular deficits nonetheless. Here we suggest that a combination of a pathogenic mutation inherited with both variants in a tri-allelic genotype may cause a large enough reduction in tyrosinase activity for a partial OCA1 phenotype.

Table 2. Predicted causal variants, in eighteen probands with phenotypes matching hypomorphic albinism. Pathogenicity determined by filtering all variants in the genes: TYR, OCA2, TYRP1, SLC45A2, SLC24A5, C10orf11 and PAX6, with the parameters MAF < 0.05, SIFT < 0.05, PolyPhen2 = possibly damaging or probably damaging. The prediction scores for non-synonymous variants are included, for some mutations a prediction score was not available at the time of analysis. Gene accessions number: TYR NM_000372, OCA2 NM_001300984, PAX6 NM_001258465, TYRP1 NM_000550, GPR143 NM_000273.
Table 3. Phenotype-genotype table of families with Sanger-confirmed TYR variants. Family number corresponds with proband number. Phenotype information (from left to right): cutaneous and hair pigmentation in context of family background, presence of nystagmus, foveal hypoplasia (FH), iris transillumination, and VEP asymmetry indicating (over)crossing of the optic nerve. Those with partial albinism are in bold.

| Family 4 | Proband | Yes - OCA1A | No | FH | No | Crossed | c.1467dup p.T489fs<sup>26</sup>,<sup>30</sup>,<sup>37</sup> | Het | Het |
|----------|----------|--------------|----|-----|-----|---------|-----------------|-----|-----|
| Father   | No       | No           | Normal | No | —   | c.1467dup p.T489fs<sup>26</sup>,<sup>30</sup>,<sup>37</sup> | WT  | WT |
| Mother   | No       | No           | Normal | No | —   | WT      | Het             | Hom |
| Sister   | No       | No           | Normal | No | —   | WT      | Het             | Het |

| Family 5 | Proband | Yes | No | FH | Yes | Abnormal | c.505<sup>-</sup>507del p.D169del<sup>44</sup> | Het | Het |
|----------|----------|-----|----|-----|-----|----------|-----------------|-----|-----|
| Mother   | No       | No  | —  | —   | —   | WT | Het | Het |
| Father   | No       | No  | Normal | No | —   | WT | Het | Het |
| Sister   | No       | No  | —  | —   | —   | WT | Het | Het |

| Family 6 | Proband | Yes | No | FH | No | Normal | c.732<sup>-</sup>733del p.C244Ter<sup>11</sup> | Het | Het |
|----------|----------|-----|----|-----|-----|---------|-----------------|-----|-----|
| Mother   | No       | No  | Normal | No | —   | WT      | Het             | Het |
| Father   | No       | No  | Normal | No | —   | WT      | Het             | Het |
| Sister   | No       | No  | —  | —   | —   | WT | Het | Het |

| Family 7 | Proband | Yes | Yes | —  | Ye s | Ye s | c.1204C > T p.R402Ter<sup>20</sup>,<sup>27</sup>,<sup>38</sup> | Het | Het |
|----------|----------|-----|-----|----|-----|-----|-----------------|-----|-----|
| Sister   | Yes | Yes | —  | —   | —   | c.1204C > T p.R402Ter<sup>20</sup>,<sup>27</sup>,<sup>38</sup> | Het | Het |
| Father   | No       | No  | —  | —   | —   | WT | Het | Het |
| Grandmother | No | No | —  | —   | WT | WT | Het | Het |

| Family 12 | Proband | No | Ye s | FH | No | Crossed | c.1217C > T p.P406L<sup>5</sup>,<sup>20</sup>,<sup>37</sup> | Het | Het |
|----------|----------|----|-----|-----|-----|---------|-----------------|-----|-----|
| Mother   | No       | No  | —  | —   | —   | WT | Het | Het |
| Grandmother | No | No | —  | —   | WT | WT | Het | Het |

| Family 18 | Proband | Yes | Ye s | FH | Mild | Inconclusive | c.1264C > T p.R422W<sup>16</sup>,<sup>39</sup> | Het | Het |
|----------|----------|-----|-----|-----|-----|-------------|-----------------|-----|-----|
| Mother   | No       | Ye s | FH | Yes | —   | —          | WT | Het | Het |

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Figure 2. Pedigree diagrams for six families with a single TYR pathogenic mutation and common polymorphism phenotyping. TYR variants are listed beneath each family. Sanger sequencing was performed on family members as opposed to the full exonic region sequenced in probands. Family number corresponds with proband number.
AROA is not an appropriate diagnosis for probands in this cohort as cutaneous and hair pigment is noticeably decreased in most probands and many family members and there is a lot of variation in ocular phenotype. Background level of pigmentation may determine the severity of the mutations as lower pigment levels will be affected more severely by the same dosage loss of tyrosinase. Therefore, our results support the theory of a causal tri-allelic genotype may go some way to account for many cases of OCA1 with apparent missing heritability. Functional studies would assist in confirming pathogenicity, thus allowing the tri-allelic genotype to be considered for both future and retrospective genetic diagnosis of OCA1.

There is potential for a double-variant haplotype, p.[S192Y;R402Q], existing on the trans allele to the known TYR mutation in affected individuals. A combination of the common variants R402Q and S192Y in cis may have a compound effect, producing a great enough loss of function equal to a deleterious TYR mutation. Each of the common variants R402Q and S192Y have a MAF of greater than 20%, and as individual SNPs they are considered benign (shown in our cohort in unaffected family members). In contrast, the predicted frequency of p.[S192Y;R402Q] in cis is 1.1%, using ‘British in England and Scotland’ participants of the 1000 Genomes project (GBR) and the webserver http://analysis-tools.nci.nih.gov/LDLink. Currently, a single variant is considered benign if the MAF is >5%56. Our findings suggest standards and guidelines could be revised to consider the combined impact of variants, particularly for more complex disorders such as albinism. Furthermore, the diagnosis of albinism currently focuses on compound mutations in single genes without considering the potential for synergistic relationships between functionally related genes such as that previously suggested for OCA2 and OCA3 genes (OCA2 and TYRP1)14 and for which there is potentially one example in our cohort.

If our proposed tri-allelic genotype hypothesis is correct, this would increase the diagnostic yield of genetic testing from 22% as described earlier, to 56% in our cohort. Given that hypomorphic albinism is a difficult cohort to diagnose clinically, evidenced by the PAX6 mutation found in the atypical case (proband 9), further exome-seq is suitable for the genetic diagnosis. A sequencing technique with broad capture allows for the pickup of genetic variants which may have resulted in an overlapping ocular phenotype.

There is no current treatment for the underlying molecular anomaly in albinism and present treatments are supportive. Therapies are under development but an effective treatment for any of the underlying molecular defects has not yet reached clinical practice. Our work and that of others appears to suggest that small variations in melanin biosynthesis between related family members dictate the extent of the phenotype in OCA pedigrees. Furthermore, the net loss of TYR function (caused by cumulative effects of multiple variants, each of which reduce TYR function by differing amounts), appear to result in a continuum of clinical features. Our work supports the assertion that small modulations in components of the melanin biosynthesis pathways, through therapeutic means, may be sufficient to rescue some of the visual disability seen in patients with albinism phenotypes.

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Author Contributions
C.S.N. and L.O.G. have contributed equally, to the wet lab and bioinformatics work in addition to manuscript preparation. They will share first authorship. J.G. and R.J.P. performed some of the bioinformatics analysis in addition to manuscript preparation and study design. D.B. and J.A.R. contributed to manuscript preparation and study design. H.G., M.R.Z., D.B., R.P., T.N., C.M. and D.W. all assisted in wet lab experiments and manuscript preparation. M.R., H.L. and F.S. all performed clinical aspects of the study and contributed to manuscript preparation. S.E. and J.E.S. contributed to study design, project oversight, bioinformatics work and manuscript preparation.

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