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Different genetic mechanisms mediate spontaneous versus UVR-induced malignant melanoma

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Abstract: Genetic variation conferring resistance and susceptibility to carcinogen-induced tumorigenesis is frequently studied in mice. We have now turned this idea to melanoma using the collaborative cross (CC), a resource of mouse strains designed to discover genes for complex diseases. We studied melanoma-prone transgenic progeny across seventy CC genetic backgrounds. We mapped a strong quantitative trait locus for rapid onset spontaneous melanoma onset to Prkdc, a gene involved in detection and repair of DNA damage. In contrast, rapid onset UVR-induced melanoma was linked to the ribosomal subunit gene Rrp15. Ribosome biogenesis was upregulated in skin shortly after UVR exposure. Mechanistically, variation in the ‘usual suspects’ by which UVR may exacerbate melanoma, defective DNA repair, melanocyte proliferation, or inflammatory cell infiltration, did not explain melanoma susceptibility or resistance across the CC. Instead, events occurring soon after exposure, such as dysregulation of ribosome function, which alters many aspects of cellular metabolism, may be important.

Introduction

Cutaneous malignant melanoma (MM) is well known to be associated with high levels of sun exposure. However, this is only true for intermittent rather than chronic exposures, with indoor workers having a higher risk for MM than outdoor workers (Gandini et al., 2005). Examples of intermittent sun exposures include number of waterside vacations and number of severe sunburns. One melanoma subtype, lentigo maligna melanoma (LMM), is invariably linked to chronic sun exposure. Individual risk for the development of MM is in part due to the number of UVR-induced mutations incurred (i.e. the environmental factor), but it is also due to genetic variation that controls skin color (degree of protection due to pigment), DNA repair capability (DiGiovanna and Kraemer, 2012), propensity to burn (inflammation), failure of programmed death of a damaged cell, and/or other factors that control melanocyte behavior (Sample and He, 2018). Genome-wide association studies (GWAS) point to genes regulating aspects of cell division control (e.g. CDKN2A/MTAP, PLA2G6, TERT), DNA repair (e.g. PARP1, APEX1, ATM), and pigmentation (e.g. MC1R, ASIP, TYR, SLC45A2, TYR) as the main players in conferring MM risk (Gerstenblith et al., 2010; Duffy et al., 2018). However only a small fraction of the variance in MM risk is explained by these genes, suggesting that there are still other genes involved in conferring MM risk in the general population (Hulur et al., 2017).

Clearly, MM is a heterogeneous disease with respect to both innate and somatic genetics, and also to environmental factors since each individual with MM was exposed to different levels of sun...
exposure. The most common superficial spreading melanomas (SSM) are sometimes found on sun-exposed body sites, but more often on non-sun exposed sites, and there can be great diversity in terms of the number of UVR signature mutations in individual lesions (Mukhopadhyay et al., 2017). While the number of UVR signature mutations in MMs strongly suggests a role for UVR in MM, the number of mutations present in a MM is due to not just obvious levels of exposure, and protective and repair differences between individuals, but also other factors, for instance the type of exposure. There is in vivo evidence that less intense exposures may be more important for skin cancer induction than more intense doses which cause more apoptosis (Lan et al., 2016), and even for a single exposure there are significant differences in skin responses between the same dose (i.e. the same levels of DNA damage) administered with high intensity over a short period versus a low intensity for a longer period (Iida et al., 2016). Our skin can protect itself via pigmentation responses (tanning), but also by ‘photo-adaption’, which is independent of pigmentation levels (Palmer et al., 2006). Thus, the skin of different individuals can respond and adapt to various forms of sun exposure in different ways, and there are potentially multiple and interacting mechanisms which might explain how UVR exposure could initiate or accelerate MM development in the general population.

For the above reasons, most experimental work on UVR carcinogenesis has used animal models. Natural genetic variation can confer resistance to many cancer types in mice (Balmain, 2002), and it is of great interest to determine why this is so. Most mouse MM models rapidly develop tumours after neonatal UVR exposure. Such models have provided tractable experimental systems to determine a MM action spectrum (De Fabo et al., 2004), to assess which type of UVR-induced DNA adducts are required, and to study the role of UVR-induced DNA damage, inflammation, and immunosuppression (Walker, 2008). It is not yet clearly known why a single neonatal UVR exposure so efficiently accelerates MM onset. We have shown previously that it is not via the acquisition of unrepaired UVR-induced damage leading to mutations in important cancer genes (Mukhopadhyay et al., 2016). A number of factors have been proposed to play a role: 1) there is a
muted inflammatory response to UVR in neonates associated with immunosuppression (Wolnicka-Glubisz et al., 2007; 15,16), Muller et al., 2008) there is a heightened sensitivity of neonatal melanocytes to proliferate following UVR (Walker et al., 209) that is driven by inflammatory cytokines, especially interferon-γ (Zaidi et al., 2011).

We used the Cdk4^{R24C}::Tyr-NRAS^{Q61K} (hereafter termed Cdk4::NRAS) mouse as a UVR-induced MM model. Somatic NRAS mutation is carried by 27% of MMs (http://www.cbioportal.org), and in ~90% of MMs the p16/CDK4/pRb pathway is deregulated via mutations in CDKN2A, CDK4, or RB1, and/or CDK4 or CCND1 amplification (Sheppard and McArthur, 2013). We studied the development of MM in mice of 70 diverse genetic backgrounds carrying these transgenes. To do so, we utilized the Collaborative Cross (CC), a set of recombinant inbred mouse strains generated from eight original founder strains, designed to enable rapid gene mapping (Churchill et al., 2004; Morahan et al., 2008). The CC is ideal for systematic analysis studies to discover modifier genes for complex diseases. Mice from each inbred CC strain may be considered as ‘clones’ of each other. Related to the CC is the diversity outbred (DO) population, in which mice descended from the same eight founders are generated as outbred stock (Churchill et al., 2012). The DO system allows very high levels of heterozygosity and recombination of CC founder alleles, but each DO mouse is genetically unique and not reproducible for experimentation requiring testing of multiple mice. The CC system has allowed us to study the influence of germline genetic background on MM induction using experimentally controlled UVR exposures. This approach tries to explain UVR-induced MM susceptibility and resistance by integrating the complex interaction of many kinds of genetic and biological information, and as such should provide much more realistic insights into MM than simple disease models focusing on single genes or proteins in isolation (e.g. Hamilton and Yu, 2012).

**Results**

**Assessment of NRAS^{Q61K} and BRAF^{V600E} transgenics as models for UVR-induced melanoma**

Before embarking on the screen for melanoma modifier genes in mice, we assessed whether there may be better murine models to work with. All models tested were on the FVB strain background. Given that BRAF^{V600E} mutation is more common than NRAS^{Q61K} in MM overall, we studied the inducible Braf^{V600E} model developed by the MacMahon lab (Dankort et al., 2009) combined with the knock-in mutant Cdk4^{R24C} mouse. Cdk4^{R24C}/R24C::Tyr-Cre^{ER}::Braf^{V600E} mice were studied in three groups. In one group, the spontaneous MM group, Braf was induced by topical tamoxifen (tam) at P1, P2, and P3. In the next group, we applied Tam at P1, P2, and P3, then exposed the mice to a single neonatal UVB dose at post-natal day 3 (P3) (Figure 1A). For the final group, we first exposed to UVR at P3, then treated with Tam at P7, P8 and P9 (Figure 1B). Surprisingly, we saw no significant difference in MM age of onset between any cohort (Figure 1C). Melanoma is not observed in Cdk4^{R24C}/R24C mice without carrying a melanocyte-specific Ras pathway mutation, with or without neonatal UVR (Hacker et al., 2006), showing that in our experiments with the Cdk4^{R24C}/R24C::Tyr-Cre^{ER}::Braf^{V600E} model, Braf^{V600E} must have been induced by the tamox application. In contrast, using the Cdk4^{R24C}/R24C::Tyr-NRAS^{Q61K} model (Ferguson et al., 2010) the single neonatal UVR exposure significantly accelerated MM age of onset (Figure 1D).

As another context in which to assess the role of the engineered mutation in mouse models of UVR-induced MM, we studied the Trp53^{ER/::Tyr-Cre(ER)}::Tyr-NRAS model in which the Trp53 deletion was induced by tamox application (26) (Figure 1E), whereas in these mice the NRAS^{Q61K} mutation is not inducible, so is present through development. Tamox treatment (i.e. Trp53 deletion in melanocytes) accelerated both spontaneous and UVR-induced MM. But there was no difference in MM onset whether or not Trp53 was deleted before or after neonatal UVR. Thus for both Braf^{V600E} and Trp53-inducible models the ability of neonatal UVR to accelerate MM may not be dependent upon whether the engineered mutation is present in melanocytes at the time of UVR exposure. Instead, it may be due to a more generalized effect via differences in DNA repair, melanocyte number and proliferative response, or inflammatory response, as has been outlined previously (Mukhopadhyay et al., 2016; Walker et al., 2009; Zaidi et al., 2011). But an oncogenic mutation in melanocytes seems to be a prerequisite. In sum, the Cdk4::NRAS model was best suited for breeding with CC mice to look for QTLs associated with UVR-dependent MM.
**QTLs for spontaneous melanoma age of onset in Cdk4::Tyr-NRAS mice**

We tested 38 CC strains to discover QTLs that modify median age of onset of spontaneous MM per strain in CC X Cdk4::Tyr-NRAS progeny (Ferguson et al., 2015). The phenotype was encoded as median age of MM onset per strain (Figure 2B), and genetic analyses performed using the Gene Miner platform (Ram and Morahan, 2017). This software uses a logistic regression matrix model over the reconstructed haplotypes matrix to produce genome-wide distribution of P values (ANOVA chi-squared). We used a false discovery rate of p<0.001 to define significant genome wide linkage. We identified a major effect QTL on mouse chromosome (chr) 16. The -Log\(_{10}\)(P) -1 interval = 14.8–21.4 megabases (Mb), a region containing 45 genes (Figure 2C). Examination of the founder haplotype coefficients in the significantly linked interval on chromosome 16 showed that the causal variant for early age of onset of MM was derived from the 129/SvJ founder (hereafter termed 129Sv). We
Figure 2. Mapping genes for spontaneous melanoma. (A) Schematic representation of breeding protocol to generate mice carrying 50% of their genome from the relevant CC strain. (B) Kaplan-Meier MM free survival curve showing age of onset on MM for all 38 strains. Strain in red shows he fastest age of onset. Note that there are four mice in the cohort, each onset value represents two mice. Blue shows slowest median age of onset. (C) Top panel shows genome-wide scan based on spontaneous MM age of onset in 38 CC strains. The average age of melanoma onset per strain was based upon onset for at least five mice for every strain. Significance of differences between groups calculated using the Log rank (Mantel Cox) test. Genotyping, construction of CC strain haplotypes, and linkage analysis was performed as described in Ram and Morahan (2017). The x-axis shows the chromosomal position and the y-axis shows the 2log10(P) values; the P-values were derived from the linkage haplotype data. Bottom panel shows plot of LOD scores along chromosome 16, with plot of the calculated log-odds ratio of eight founder alleles over the chromosome where the founders are color-coded. (D) Genes containing potentially regulatory intronic, 5’ or 3’ UTR variants. (E) Expression level of genes in the HAIR-GEL skin gene expression database (28). Y axis denotes FPKM (Fragments Per Kilobase Million). (F) Gene expression fold changes in mouse skin from 129S (which carries the susceptibility allele) compared to C57BL/6, NOD, and FVB, which do not. Based on gene expression values from RNA sequencing. Lamp3 is separated from the other genes since it is the only one in the figure not located near a 129S-specific regulatory SNP defined in the ENCODE database.

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ascertained which genes within the interval were the best candidates by cataloguing DNA variants that are carried on the causal 129S haplotype, and not any of the other founder haplotypes. Mining the Sanger Mouse Genomes and ENCODE databases for 129S-specific variants revealed eight candidates, two of which (Prkdc and Arvcf) carry non-synonymous mutations (Figure 2D), while all 8 (Prkdc, Pkp2, Yars2, Arvcf, Gp1bb, and Abcc5) had 129S-specific single nucleotide potentially regulatory polymorphisms (SNPs) in their 5' or 3' UTR, or introns. It is possible that one of the 129S-specific SNPs near the candidates may regulate another gene elsewhere in the genome, or that there are other 129S-specific regulatory SNPs nearby any of the 45 genes in the region that were not detected by ENCODE.

Next, we looked at skin gene expression of the six candidates. They are expressed in neonatal mouse skin and/or hair follicle, including in melanocytes, as published in the Hair Gel database (Sennett et al., 2015) (Figure 2E). We then performed global gene expression analysis (RNAseq) (Supplementary file 1) of skin from adult mice from various laboratory strains (AJ, NOD, B6, FVB, and 129S) (Figure 2F), four of which are CC founders, for which the genome sequences are available in the Sanger database. We found no differences between susceptible (129S allele-carrying) and resistant strains (non 129S allele-carrying; AJ, NOD, FVB, B6) that would help demarcate a candidate(s). Only one gene (Lamp3, lysosomal-associated membrane protein 3) of the potential expressed sequences within the region was significantly differentially expressed between the susceptible and resistant strains assessed (Figure 2F), but the nearest 129S-specific regulatory SNP is located 10 Kb away, near Gp1bb, and the next about 500 Kb away, near Avcrf. Unfortunately, there is no skin expression quantitative trait locus (eQTL) data for mouse skin, and it was not possible to consult the GTEx eQTL database for eQTLs in human skin since the LAMP3, GP1BB, and ACVRF genes in humans are scattered on different chromosomes.

Therefore, we hypothesized that a functional effect on phenotype was most likely the presence of missense mutations rather than regulatory expression changes on the causal allele. Prkdc carried two missense mutations, one defined as deleterious by SIFT. This is the same mutation previously mapped and validated as a modifier of lymphomagenesis (missense mutations rather than regulatory expression changes on the causal allele. Genes in humans are scattered on different chromosomes. We consulted the GTEX eQTL database for eQTLs in human skin since the is no skin expression quantitative trait locus (eQTL) data for mouse skin, and it was not possible to determine whether variation in its expression is correlated with expression of these GWAS breaks (e.g., PARP1, ATM, APEX1) are in linkage disequilibrium with SNPs associated with melanoma risk in GWAS (Hulur et al., 2017). We reasoned that even though PRKDC is not a GWAS hit, we could determine whether variation in its expression is correlated with expression of these GWAS genes. We examined global gene expression in non-sun exposed human skin across the GTEx cohort (Genenetwork.org). Networks of the top 500 genes correlated with PRKDC were constructed at a confidence value of 0.9 using STRING (https://string-db.org). The most significant network for molecular function was RNA binding (p=1.22 ×10^-37 false discovery rate), and in KEGG pathways DNA replication (p=1.22 ×10^-8). PARP1 (at 170, r = 0.46, p=2×10^-16), and APEX1 (at 287, r = 0.43, p=8×10^-14) were in the top 300 most significantly correlated genes with PRKDC. ATM was not in the top 500, but its relative ATR was at number 105 (r = 0.49, p=4×10^-16). While these correlations are based only on gene expression across the GTEx cohort, not any other aspect of gene function, they do point to the possibility that PRKDC is associated with pathways associated with DNA double strand break repair, components of which are encoded by other genes which confer MM risk in the general population.
QTLs for UVR-induced melanoma age of onset in Cdk4::Tyr-NRAS mice

CC-transgenic progeny strains from 70 CC strains were exposed to a single neonatal exposure then followed until MM developed (Figure 3A). Median age of melanoma onset per strain was scored as the phenotype (Figure 3B). Only lesions developing on the UVR-exposed dorsal surface (where the overwhelming majority developed) were counted. We identified a major effect QTL on mouse chr.1 (-Log_{10}(P) – 1 interval = 187.8–189.2 Mb) (Figure 3C), a region containing 10 genes. We ascertained which genes within the interval were the best candidates by cataloguing DNA variants on the causal allele that vary between susceptible or resistant strains. The causal allele was carried by AJ and NOD. Mining the Sanger Mouse Genomes and ENCODE databases for NOD/AJ-specific variants revealed four candidates carrying variants specific to the causal allele (Tgfb2, Rrp15, Spata17, and Gpatch2). Rrp15 was the only one carrying missense mutations (Figure 3D), while the other three genes (Tgfb2, Spata17, and Gpatch2), had AJ/NOD-specific single nucleotide polymorphisms (SNPs) in their 5’ or 3’ UTR, or introns (Figure 3E).

Next we looked at the Hair Gel database to determine whether these genes were expressed in skin. All but Spata17 were, essentially ruling this gene out as a candidate (Figure 3F). We then looked at skin gene expression (Supplementary file 1) between AJ, NOD, and FVB, which carry the Chr. one allele, and B6, DBA, and 129S which do not (Figure 3G). There were no significantly differentially expressed genes between groups, suggesting a missense rather than a regulatory causal variant. Thus, while we cannot rule out Tgfb2 and Gpatch2, by those criteria Rrp15 is the best candidate. The causal allele (AJ/NOD) carried two Rrp15 missense variants, both defined by SIFT as likely to be ‘tolerated’. But one at amino acid 117 is most likely to be the causal mutation given that it is a Glu > Gln change, whereas the other is Ala > Val, which is likely to be silent. Rrp15 encodes a ribosomal subunit that is part of pre 40S and pre 60S subunits that is important for rRNA transcription and ribosome biogenesis. Furthermore, Rrp15 knockout in vitro causes nucleolar stress by activating the Mdm2-Trp53 axis, and subsequently a G1-S phase cell cycle blockage (De Marchis et al., 2005; Dong et al., 2017). Deregression of the ribosome complex decreases the fidelity and patterns of mRNA translation and many other downstream events (Quin et al., 2014; Pelletier et al., 2018). Although much of the effect on cell behaviour is via altering the stabilisation of p53 by Mdm2, this can also occur via p53-independent mechanisms (James et al., 2014).

Confirmation of the chromosome one locus using diversity outbred (DO) mice

DO mice harbor frequent recombinations throughout their genomes. We hypothesized that some DO animals (Churchill et al., 2012) would have informative recombinants of the causal AJ/NOD allele to make them useful for fine mapping the linked chr.1 interval. We tested DNA from 314 DO mice, initially using six polymorphic markers across an 8 Mb region around Rrp15. We selected a subgroup of 8 DO mice with apparent recombinations within the AJ/NOD alleles across the region and genotyped them using a panel of 60 SNPs (20 of which are the most informative and shown in Supplementary file 2). SNPs were chosen based on their ability to discriminate the AJ/NOD allele from the other eight founders, especially those variants private for AJ, NOD, or both. While this NOD/AJ allele was scattered throughout the region in different DO mice, as expected we observed many recombinations across the chr.1 region of interest. However there were many gaps and alleles for which we could not unambiguously call the founder haplotypes on both strands across the region (Supplementary file 2). To adequately do this, many more SNPs would have to be tested. Nonetheless we crossed each of the eight selected DO mice with Cdk4::NRAS transgenics, and studied UVR-induced MM onset in the progeny (Figure 3H). Because of the limitations of our SNP-based map of the region, to genotype the DO mice we used manual Sanger sequencing to genotype more densely, in particular in the region of our candidate gene Rrp15 (Figure 3I). As seen in Figure 3H, the predicted causal SNPs in Rrp15 do not segregate with fast MM onset as they did in the CC strains. We hypothesized that lack of penetrance of the Rrp15 allele was due to the introduction of additional resistance alleles elsewhere in the genome due to the high levels of recombinations and heterozygosity in the DO genomes. We tried to circumvent this by backcrossing transgenic-DO mice onto C57BL/6. Mice from several litters were followed, and after two backcrosses we assessed UVR-induced MM age of onset, with progeny genotyped immediately around Rrp15 by Sanger sequencing. We found that the penetrance of the causal Rrp15 variant was restored on backcrossing: mice
Figure 3. Mapping genes for UVR-induced melanoma. (A) Schematic representation of breeding protocol to generate mice CC progeny. Pups were exposed to a single UVR exposure at post-natal day 3. (B) Kaplan-Meier MM free survival curve showing age of onset on MM for all 38 strains. Strain in red shows the fastest age of onset. There are two data points (four mice) for this strain cohort, but each onset value represents two mice that developed MM at the same time. Blue shows slowest median age of onset. (C) Top panel shows genome-wide scan based on neonatal UVR-induced MM onset (days). Bottom panel shows QQ plot for genome-wide scan. (D) Candidate gene expression in neonatal mouse skin (www.hair-gel.net). (E) Fold change in mouse skin susceptible (AJ/NOD allele) vs resistant (non-AJ/NOD) background. (F) Effectiveness of UV in MM (days). (G) Top panel shows genome-wide scan based on neonatal UVR-induced melanoma. Bottom panel shows QQ plot for genome-wide scan. (H) Spontaneous MM and UVR-induced MM progeny. (I) Effectiveness of UV in MM (days). (J) Cdk4:Nras::D07 X C57BL/6 backcross progeny. (K) Days to MM onset. (L) Albinism strains vs pigmented strains. Figure 3 continued on next page.

Ferguson et al. eLife 2019;8:e42424. DOI: https://doi.org/10.7554/eLife.42424
carrying the NOD/AJ alleles of Rrp15 had significantly earlier MM onset (Figure 3J). Thus, the propensity for neonatal UVR to accelerate melanoma is highly dependent upon genetic background influences. In addition, there are additional potential resistance alleles which can, if present, interact with the Rrp15 susceptibility allele.

**Neonatal UVR acceleration of melanoma age of onset is dependent upon innate genetics**

We further analyzed the role of neonatal UVR in accelerating melanoma age of onset by subtracting the average age of onset of spontaneous from that of UVR-accelerated melanoma (Figure 3K). This provides us with a further phenotype: the strain-specific rate of acceleration of MM onset by neonatal UVR (Figure 3L). The median age-of-onset for UVR-induced MM per strain was subtracted from the median onset for spontaneous MM, to denote what we have termed the ‘effectiveness’ of UVR in inducing MM. In resistant CC strains, neonatal UVR does not accelerate MM onset at all (e.g., HOE, BAX2), or does so by a very small amount, whereas for susceptible strains MM onset was accelerated by more than 300 days (e.g., SEH, XAJ2) as seen in Figure 3K and L. The phenotype is independent of pigmentation status (albino vs pigmented) (Figure 3L). The genome scan for this trait is shown in Figure 3M. Unfortunately, we only have both spontaneous and UVR-induced onset data for 27 strains. While this does not provide enough power to detect genome-wide significant linkage to this phenotype, there was a peak of suggestive significance at chromosome 1 p, overlapping with the QTL detected for UVR-induced age of onset, containing the Rrp15 gene, with the NOD allele showing a different coefficient from the other strains.

**Skin gene expression changes after neonatal UVR**

The way by which neonatal UVR accelerates MM may provide insights into early events in the initiation of this neoplasm. To search for pathways deregulated in neonatal UVR-exposed skin, we performed global gene expression chip studies with Illumina Beadchips on UVR exposed and non-exposed epidermis of wild-type FVB mice at various time-points (Supplementary file 3). The top 500 significantly up- and down-regulated genes after neonatal UVR were used to construct networks for UVR-induced gene expression changes at a confidence value of 0.9 using the STRING resource (https://string-db.org) (Figure 4). At 6 hr after UVR exposure, one can see strong significant evidence of ribosome biogenesis occurring, with downregulation of various metabolic pathways compared to control untreated skin (Supplementary file 4). At 10 hr, post-UVR metabolic activity is still suppressed, but DNA replication is either already occurring to some extent, or about to occur for repair and reconstruction of damaged cells. Upregulated p53 signaling was also observed. At 24 hr post UVR various pathways involved in cell proliferation (ribosomes and translation, metabolism) and
reconstruction (extracellular matrix, metabolism) are significantly activated in the epidermis. One of the problems with using whole tissue is that one cannot discriminate in terms of cell type. As a way to put the skin gene expression into some perspective we wanted to look at skin gene expression changes at some days after UVR, since it is known that this period corresponds with an influx of immune cells into the dermis (Zaidi et al., 2011) and the influx of melanocytes into the epidermis (Walker et al., 2009). Therefore, we separately studied gene expression in the neonatal epidermis and dermis harvested at 3d after UVR (Figure 5A). There was a striking loss of immune markers in the epidermis, mostly reflecting UVR-induced migration of epidermal Langerhans cells to the dermis as expected. We also saw a strong gene expression cluster for genes involved in melanogenesis, reflecting the presence of melanocytes in the UVR-exposed epidermis, but not in control skin at the time-point 3 d after UVR. In the dermis, we observed networks reflecting increased cellular activity, in particular upregulation of kinases and interferon-induced enzymes. But in contrast to the epidermis, in the dermis we also observed a signal for myeloid cells, presumably reflecting the influx of macrophages, which occurs maximally at around this time post UVR (Handoko et al., 2013).

The critical question with respect to genes deregulated by neonatal UVR is the behavior of our candidate genes, one of which must be causal in accelerating MM. Rrp15 was the only one of the chr.1 candidates significantly changed in expression by UVR (Figure 5B). Hence in addition to the
Figure 5. Gene expression networks induced by UVR. (A) Gene network analysis showing at epidermis (top panel), and dermis (bottom panel) 72 hr after neonatal UVR, the top 500 genes (all p<0.05) deregulated in UVR-treated neonatal epidermis versus untreated (from the same mouse on the side blocked with electrical tape). Pathways labelled in red were upregulated after UVR, and those in blue downregulated. For those labelled in black some genes were up and some downregulated. (B) Fold change of expression of candidate genes after neonatal UVR. The only gene within the candidate Figure 5 continued on next page
sustainable CC strains carrying a missense mutation in the putative Rrp15 causal allele, the fact that it is the only candidate that responds to neonatal UVR adds weight to it being the best candidate in the linked genomic region. Our gene expression analysis was performed on whole skin, epidermis, or dermis, rather than individual cell types, allowing signals from a number of cell types, for example keratinocytes, immune cells, and melanocytes, to be taken into account. Since Rrp15 is expressed virtually ubiquitously in the skin we do not know whether its mode of action in accelerating UVR-induced melanoma would be cell intrinsic or extrinsic. But we note that in a study of UVR-treated cultured melanocytes, ribosome metabolism was the most significantly altered pathway at 6, 12, and 24 hr after UVR (López et al., 2015), as in our whole skin studies. In addition, to gain a better sense of changes in skin correlated with RRP15 expression we constructed gene networks based on the top 500 genes correlated with RRP15 expression across the sun-exposed anatomical site human skin GTEx cohort (Figure 5B). As expected changes in RRP15 gene expression are correlated with ribosome biogenesis, but also many other aspects of cell behavior, including DNA damage recognition and repair, transcription, and splicing. Defects in any of these processes could in theory explain why MM is accelerated by UVR exposure not only in mouse strains which carry germline Rrp15 variants, and putatively in humans also.

Mechanism by which melanoma is accelerated by UVR
Since genetic background greatly influenced whether or not MM was accelerated by neonatal UVR, we studied various effects of UVR a few days after exposure on traits that may differ between susceptible and resistant strains. These included the rate of removal of UVR-induced cyclobutane pyrimidine dimers (CPDs), influx of inflammatory neutrophils and macrophages, and the proliferation of melanocytes (increase in melanocyte number in the epidermis) (Figure 6A). We again used readily-available laboratory strains (AJ, B6, DBA, FVB, NOD, 129 s), four of which are CC founders. For all six, we determined the median age of onset of MM after neonatal UVR. We chose time-points 1, 4, and 7 days post UVR. In C57BL/6 mice, CPDs are generally removed by d4 after UVR, certainly by d7. Neutrophils generally infiltrate the skin by d1 after UVR and numbers decrease by d4. Macrophages are maximally present at 4d (39). At 4d, melanocytes have migrated to the epidermis in response to UVR, macrophages are at their maximum number, roughly concurrent with maximum epidermal melanocyte density. By the d7 post UVR time-point, the skin is ostensibly returned to normal, but in some mouse strains there may be a delay in some measures.

First, across the six laboratory strains we found no differences in the capacity to remove CPDs, as measured by the proportion of skin cells carrying them post-UVR. However, there was great variation between the various strains in the size and timing of the melanocyte and myeloid cell responses after UVR (Figure 6A). To ascertain whether the level of any of these responses could be correlated with age of MM onset, we compared results for the lab strains stratified in terms of whether they carry the susceptible or resistant allele at chr.1 (e.g. around Rrp15) (Figure 6B). When compared in this way none of these commonly purported mechanisms by which UVR exacerbates MM were significantly associated with onset of UVR-induced MM, although we found a non-significant trend for the susceptible strains to show higher levels of neutrophil influx at d1 after UVR. As an additional assessment, we performed a correlation test between MM age of onset and the phenotypes measured and found no significant correlation, although d1 neutrophil infiltration was nearest to significance.

If one accepts that the low number of strains compared (3 vs 3) makes it difficult to attain statistical significance for a mechanism with a complex milieu of events, one could argue that neutrophil influx may have an effect on exacerbating MM. Thus we set out to determine whether depletion of neutrophil infiltration using a neutrophil-blocking anti-Ly6g antibody would influence subsequent melanoma induction in Cdk4::NRAS mice. We injected blocking antibody at 1d before UVR, the day of UVR, and 2d after UVR. First we assessed neutrophil depletion after three injections by taking skin...
Figure 6. Immune and melanocyte proliferative responses post UVR in different mouse strains. (A) Parameters measured post UVR in different mouse strains. Top bar graph shows number of cells (stained with anti-CPD antibody) positive for pyrimidine dimers at 1 (D1), 4 (D4) and 7 (D7) in whole skin after neonatal UVR. Second bar graph shows the number of epidermal melanocytes (stained with anti-Sox10 antibody) per field. Third graph shows the number of dermal neutrophils (stained with anti-myeloperoxidase) per field. Fourth graph shows the number of dermal macrophages (stained with anti- Figure 6 continued on next page

Ferguson et al. eLife 2019;8:e42424. DOI: https://doi.org/10.7554/eLife.42424

Cancer Biology
sections and staining with neutrophil-specific anti-myeloperoxidase antibody (Figure 6C). Anti-Ly6g treatment significantly depleted neutrophil influx after neonatal UVR (Figure 6D). We exposed two cohorts of Cdk4::NRAS mice to anti-Ly6g anti-neutrophil depleting Ab, or PBS control respectively, then performed a longitudinal study for MM age of onset, but saw no difference in the age of onset of MM between cohorts (Figure 6E). Thus, inhibition of the neutrophil influx did not suppress UVR-induced genesis.

Discussion

There are likely to be multiple and interacting mechanisms which might explain how UVR exposure could initiate or accelerate different types of MM. We have used a mouse MM model to study the development of MM on many CC strain backgrounds. The Cdk4::NRAS model we used is a well-characterized model of UVR-induced MM, and in terms of an acceleration of MM onset after neonatal UVR it behaves similarly to other models which carry constitutive oncogenic mutations in melanocytes (e.g. the Mt-Hgf model) (Noonan et al., 2001). But, surprisingly, we found that MM was not accelerated in a model carrying an inducible BrafV600E mutations in its melanocytes. It is known that modes of engineered oncogene expression in melanocytes can influence other phenotypes also. For example, BrafV600E induced in melanocytes in utero is incompatible with embryonic viability (e.g. Dankort et al., 2009), whereas in a non-inducible transgenic BrafV600E model it is not (Wurm et al., 2012). We do not know why MM was not accelerated in the inducible BrafV600E model, but one possibility is that there may be simply more activated melanocytes in the neonatal Tyr-NRASQ61K transgenics at the time of UVR exposure since NRASQ61K is expressed in utero, whereas BrafV600E is induced only for a short period of 3 days before UVR (i.e. P1-P3). We favour such an explanation rather than a mechanistic difference between the respective oncogenes (Braf vs NRAS) per se, since in another similar inducible BrafV600E model repeated UVR exposures significantly accelerate MM when the oncogene is activated at 4 weeks of age (Goel et al., 2009). In a model using inducible Trp53 deletion instead of constitutive Cdk4R24C we found that UVR accelerated MM, but whether Trp53 was deleted before or after UVR was inconsequential. This could be somewhat akin to the work of the Evans laboratory who showed that although TP53 is very important in helping modulate the acute DNA damage response to radiation, its function in this period does not contribute to protection against transformation, in fact it is only within the long lag phase leading to cancer development that TP53 acts as a tumour suppressor (Christophorou et al., 2006). Therefore, mouse models appear to differ with respect to whether and by how much neonatal UVR accelerates MM. There is no perfect animal model for ‘generalized’ MM, but one would expect that the Cdk4::NRAS transgenic in combination with the CC should provide useful information as to how genetic background can influence UVR-induced murine MM.

Age of onset of spontaneous MM across the CC was underpinned by germline variation in the Trp53 gene on chr.16. This is overwhelmingly the best candidate in the mapped interval as the mutation carried on the 129S susceptibility allele has been functionally validated in other mouse models.
of cancer (Yu et al., 2001; Degg et al., 2003; Goodwin et al., 2015). Although it is known to be involved in recognition and removal of DNA damage, it is unclear whether its function in MM would be cell intrinsic or extrinsic, since it is ubiquitously expressed in skin cell types. In addition, this is quantitative genetics, and although the QTL explains a large proportion of the phenotypic variation, other cooperating loci may also exist, for instance on chr.5, (Figure 3C), which might become statistically significant (or disappear) if more CC strains were tested. In terms of possible human relevance, other genes (e.g. PARP1, APEX1) involved similarly in DNA repair, and whose expression is correlated with that of PRKDC in human skin, are associated with population-based MM risk (Duffy et al., 2018). Thus PRKDC may be an example of a modifier of MM in mice, but mediated by other genes of the same pathway in humans. But the mechanistic result of Prkdc deregulation might be similar to the effects of deregulating MM risk genes like PARP1 and APEX1, that is, all ultimately influencing DNA repair. Hence our genetic screen appears to have revealed mechanistically relevant information, notwithstanding that the particular causal gene variants can be different between mouse and man. One can also consider Prkdc as a MM modifier gene, that is, our genetic screen detects MM modifiers in the context of a mouse model which carries a germline Cdk4 variant. In keeping with this, PRKDC was also one of the few DNA repair genes associated that modified MM risk in MM-prone families (Liang et al., 2012), many of which carry a CDKN2A mutation, and a few CDK4 mutation. If we assume that Cdk4::NRAS mice may in some respects be ostensibly a model for familial MM, the findings of Liang et al., 2012 further support the possible relevance of our findings in human disease.

One of the most striking findings from our study was that the genetic polymorphisms modifying MM onset were very different between spontaneous and UV-induced disease (Figure 7). This is particularly notable since in this model MM is accelerated by just a single UVR exposure. The age of onset of neonatal UVR-induced MM in the transgenic-CC progeny across the CC was linked to a chr.1 locus containing a strong candidate, Rrp15, with a missense mutation. Taken together with the fact that of the candidates within the linked region, only Rrp15 was significantly differentially expressed between susceptible and resistant strains, only Rrp15 was upregulated in the epidermis of
neonatal skin after UVR exposure (at 24 hr), we deemed it the strongest candidate. In addition, gene ontology and KEGG pathway analysis of UVR-induced gene expression changes in neonatal mouse skin revealed that ribosome biogenesis is one of the major gene networks upregulated at 6 and 24 hr after UVR. One caveat here is that for the UVR-induced gene expression studies we used whole epidermis (or dermis). Notwithstanding that there are differences in melanocyte and immune cell density over time between 1 and 3 d after neonatal UVR, most of the signal would have come from keratinocytes at all time-points since they are the most numerous cells in the epidermis. But our analysis of UVR-induced gene expression changes, based upon fold-change, are sensitive enough to pick up gene networks that reflect predictable changes in melanocyte and immune cell density (although not the type of immune cells). Rrp15 is expressed in all cell types within the neonatal skin (Figure 3F), but we do not know whether its mode of action in modifying melanocyte transformation is cell autonomous or non-cell autonomous. We assume that the upregulation of ribosomal metabolism we observe at 24 hr after UVR reflects mostly changes in keratinocytes, but very similar responses have also been found in UVR-treated cultured melanocytes (López et al., 2015). Single cell sequencing of different skin cells may help resolve these problems, but even such methods also have their drawbacks including unwanted gene expression changes induced soon after removal of cells from their microenvironment.

We have used inferences taken from a variety of sources, which are by themselves not confirmatory, to build a strong case for a role for variation in Rrp15 function influencing the propensity for UVR to accelerate MM. There is some evidence that SNPs near RRP15 are also associated with human MM in some contexts. The world melanoma genetics consortium found such evidence using a method for candidate genetic association to detect variants that may not reach genome-wide statistical significance after correction for multiple testing (Schoof et al., 2012; Wurm et al., 2012). There was no information on individual sun exposure in the tested cohorts. Of the 39 immune-related genes tested, SNPs near LGAL3 and TGFB2 were the most significantly associated with MM risk (Rendleman et al., 2013). Of note, the RRP15 and TGFB2 genes are located adjacent to one another and just 7 kb apart. Which of the two is the causal gene is difficult to elucidate in the human genetics study, but our systems analysis work using the CC has allowed us to build a very strong case for Rrp15.

We discovered that genetic background dramatically influences the propensity for melanocyte transformation after UVR. As well as discovering a gene likely to be associated with this, we also examined differences in skin responses between susceptible versus resistant mouse strains. We did not observe major differences in the propensity for removal of UVR-induced CPDs, nor in melanocyte proliferation, nor macrophage influx. We observed a weak correlation with the number of skin-infiltrating neutrophils at d1 after neonatal UVR and UVR-induced MM onset. However, depleting neutrophils before and after neonatal UVR did not reduce the time of onset of MM, somewhat in line with our previous finding that depletion of macrophage infiltration also did not abrogate the MM-accelerating effect of UVR (Handoko et al., 2015). It could be argued that by using immunohistochemical staining to assess removal of UVR-induced CPDs we could miss subtle differences between mouse strains in the repair process that may be consequential in terms of leaving a molecular memory of UVR damage in the transformed cells. But even our exome sequencing on the neonatal UVR-induced mouse melanomas does not suggest a major role for UVR-induced mutagenesis (Mukhopadhyay et al., 2016).

Researchers tend to look for measurable skin responses to UVR exposure to try to understand how melanocytes may be destabilized and ultimately transformed by DNA damage. The ‘usual suspects’ include defective DNA repair, photo-immunosuppression, inflammation, and cell proliferation, which are observable in the days following exposure. These can all enhance carcinogenesis in specific contexts. But our results suggest that earlier events in the few hours post UVR such as aberrant ribosome activity that can cause inappropriate protein expression, may be more important. It is not clear whether this would be acting during the acute damage repair period after neonatal UVR, or during the lag period leading to tumour initiation. The notion of a particular constellation of gene networks that vary between CC strains and confer resistance to the MM-accelerating effects of UVR may not be dissimilar to what occurs in amphibians, where regenerating limbs, but not non-regenerating body parts, are resistant to carcinogen-induced cancer (Sarig and Tzahor, 2017), despite the fact that both anatomical sites incur the requisite DNA damage. Particular molecular networks within skin cells in the MM-resistant strains appear to work against transformation, despite animals from all
strains being exposed to high levels of UVR-induced damage. Hence in keeping with the fact that even non-cancerous human skin can carry UVR signature mutations in cancer genes (Martincorena et al., 2015), incorrectly repaired UVR-induced DNA damage leading to somatic mutation is in essence necessary but not sufficient to exacerbate MM, and the presence of germline variants for melanoma susceptibility and resistance is very important. Biological systems (e.g. skin and skin UVR responses) are very complex and varied across a population of different individuals. Our systems analysis strategy has attempted to harness such complexity and in doing so has resulted in the discovery of some potentially important findings with regards UVR-induced melanoma. We have performed a genetic screen for natural genes regulating MM age of onset and found surprisingly that different genes mediate spontaneous and UVR-induced MM susceptibility (Figure 7). We have identified a strong candidate genes and potential mechanisms in both cases.

Materials and methods

Key resources table

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### Mouse melanoma model

*Cdk4^{R24C/R24C}::Tyr-NRAS^{Q61K/+}+* mice are previously described in *Ferguson et al. (2010)*. We crossed mice with breeding partners from each CC strain (*Ferguson et al., 2015*). Hence all study mice are *Cdk4^{R24C/+}::Tyr-NRAS^{Q61K/+}.* All experiments were undertaken with institute animal ethics approval (A98004M). Mice were sacrificed before tumors exceeded 10 mm in diameter. In some melanoma-resistant strains lymphomas developed in some mice at >400 days of age. Such mice were counted as melanoma-free at the age of death. Each phenotypic measurement is based upon at least 4 and up to 15 mice per CC strain background. *p53^{F/F}+* mice (carrying floxed alleles allowing Cre-mediated excision of exons 1–10) were obtained from the Mouse Models for Cancer Consortium (*http://mouse.ncifcrf.gov*). Melanocyte-specific Trp53 deletion in *p53^{F/F}+/Tyr-Cre (ER)/Tyr-NRAS* mice was induced via topical application of 8-OH-tamoxifen (15 mg/ml in DMSO) at P0, 1 and 2. For the studies involving the BRAF model we used the inducible BRAF^{V600E} model generated by (24). All these mice were of FVB strain background.

### CC breeding

Collaborative cross mice were obtained from the Animal Research Council (ARC) in collaboration with Prof Grant Morahan of the Harry Perkins Institute of Medical Research, Perth, Australia. A/J, C57BL/6j (B6), 129/SvJ (129S), DBA, and NOD/ShiLtJ (NOD) were purchased from the Animal Resources Centre, Western Australia.

### UVR treatments

Pups (3-day-old) were exposed to a single UVB exposure from a bank of 6 Phillips TL100W 12RS UVB lamps (Total UVB dose, 5.9 kJ/m², or an erythemally-weighted dose of 1.8 kJ/m²) UVB.

### Phenotypic characterization

We have previously described a system for visual tracking of lesions developing on the FVB *Cdk4^{R24C/R24C}::Tyr-NRAS^{Q61K/+}+* mice and a histology-based staging system (*Wurm et al., 2012*) and mice on the various CC strain backgrounds were scored in this way. Briefly, lesions were excised after death followed by conventional histopathologic work-up with haematoxylin and eosin (H and E) staining. Each lesion was viewed and confirmed individually as MM by BF and GJW as described in Wurm et al. (42). Where there was any doubt about diagnosis, tumors were stained with Trp1 and/or Sox10. No skin tumors were observed apart from melanomas. We included in our analyses...
melanomas that developed on dorsal surface only. Mice were sacrificed before tumors exceeded 10 mm in diameter.

**QTL analysis**
The construction of the CC founder haplotypes is described in Ram and Morahan (2017). For mapping we used a logistic regression matrix model over the reconstructed haplotypes matrix to produce genome-wide distribution of P values (ANOVA chi-squared). We used a false discovery rate of p=0.0001 to define significant genome wide linkage.

**SNP genotyping**
We used a custom SNP approach performed by AGRF, a custom array of 92 SNPs across the 180–190 Mb region of mouse Chr. 1. Genotyping was performed using the Sequenom mass array system.

**Sanger sequencing**
Target fragments chosen to contain multiple SNPs were PCR-amplified, then cleaned from excess primers and nucleotides using CleanSweep PCR Purification (Applied Biosystems, Life Technologies, Carlsbad, CA, USA). Sequencing was carried out using BigDye Terminator v3.1 Cycle Sequencing and then run on ABI Prism DNA Sequencers (Applied Biosystems). The sequencing traces were compared using Multiple SeqDoc chromatogram comparison programme (http://research.imb.uq.edu.au/seqdoc/multi.html).

**Mouse expression array**
Five hundred ng of total RNA from each tumor was used as the starting material to produce cRNA, following Illumina TotalPrep RNA Amplification protocol. From each sample, 1500 ng of cRNA were hybridised to Illumina MouseWG-6 v2.0 Expression BeadChips (Illumina, San Diego, CA, USA) and then scanned. Data were extracted using Genome Studio (Illumina, San Diego, CA, USA) and then imported into GeneSpring GX 11.5.1 (Agilent Technologies, Santa Clara, CA, USA), before subsequent analysis. Normalisation was performed using the R package LUMI, and differential analysis done using LIMMA. Multiple testing correction was carried out using the Benjamini-Hochberg procedure. Differentially expressed genes with p-adjusted values < 0.05 were considered significant.

**RNA seq**
Skin gene expression experiments were undertaken with institute animal ethics approval (A98004M). RNA was isolated using RNeasy Kit and libraries generated with Illumina mRNA kit. Sequencing was performed using Illumina HiSeq chemistry, with 50 bp single reads. RNA-seq samples were mapped to mouse genome MM10 using STAR. Quality control metrics were computed using RNA_SeQC version 1.1.18, and expression values estimated using RSEM version 1.2.30. We corrected for library size by dividing each sample’s count by millions of reads mapped. We used the ‘calcnormfactors’ function from the EdgeR package to obtain TMM factors and used these to correct for differences in RNA composition.

**Gene expression correlation and gene network study**
To generate interconnected networks based on correlations, gene lists were clustered using STRING (http://string-db.org/). STRING creates networks representing the best available knowledge of gene interconnections. Each protein-protein interaction is annotated with ‘scores’ indicating how likely an interaction should be true. Scores rank from 0 to 1, with one being the highest confidence. A score of 0.5 indicates roughly every second interaction might be erroneous. Gene-gene co-expression correlations were computed as Pearson product-moment correlations (r) in Genenetwork.org after removing outliers.

**Antibody staining**
Skin from the pups at Day one after UVB radiation (D1), D4 and D7 was paraffin embedded. All immunohistochemistry staining was performed on pup skin sections (4 um) with standard DAB or NovaRed. Counts were performed on multiple fields from multiple skin sections from each of >3...
mice (each field is ~1 mm in length). Macrophage staining used F4/80 from CD68 rat monoclonal antibody diluted to 1:400, Abcam ab6640, Cl: A3-1 (Cambridge, UK). Melanocyte nuclear staining used Sox10 from sc-17342 goat polyclonal antibody (N-20) diluted to 1:200, Santa Cruz Biotechnology (Dallas, TX, USA). Cyclobutane pyrimidine dimers (CPD) staining was with monoclonal anti-thymine dimer, Clone H3 (T1192) diluted to 1:400, Sigma-Aldrich (St Louis, MO, USA). For this CPD staining, sections were first blocked in 1% hydrogen peroxide, then incubated in 50% ethanol, 30% ethanol/0.02N HCl, 0.05N HCl, and 0.07N NaOH/70% ethanol, before incubating in primary antibody. Neutrophils were stained using an anti-Ly6G as primary antibody, except after neutrophil depletion, when staining was done using MPO staining (see below). Anti-Ly6G was from rat anti-neutrophil monoclonal antibody Abcam ab2557: NIMP-R14 (Cambridge, UK) at a concentration of 1:100.

**Neutrophil depletion**

Each NOD/Cdk4^R24C+/::NRAS pup was injected intraperitoneally, with either 100 ug InVivo MAAb anti-mouse Ly6G (Bio C Cell, Beverly, MA, USA) or PBS (as a control) at P2, P3 and P5. At P3, all pups were also given UVR treatment. As the depletion was done using anti-Ly6G, staining to differentiate between depleted and non-depleted skin was done using Myeloperoxidase staining (Abnova rabbit anti-MPO, Taipei, Taiwan) at a concentration of 1:75.

**Statistical analysis**

One-way ANOVA tests were used to determine the significant difference between means, using R. The survival of mice in each treatment group was estimated using Kaplan-Meier analysis (PRISM™), and the Log-Rank (Mantel-Cox) test was used to test for differences between the groups. For correlation comparisons Pearson correlation r value was calculated in PRISM along with the p-value for significant correlation.

**Acknowledgements**

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**Additional information**

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The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

**Author contributions**

Blake Ferguson, Herlina Y Handoko, Conceptualization, Data curation, Formal analysis, Validation, Investigation, Methodology, Writing—review and editing; Pamela Mukhopadhyay, Data curation, Software, Formal analysis, Methodology, Writing—review and editing; Arash Chitsazan, Data curation, Investigation; Lois Balmer, Resources, Data curation, Methodology; Grant Morahan, Conceptualization, Resources, Formal analysis, Supervision, Funding acquisition, Methodology, Writing—review and editing; Graeme J Walker, Conceptualization, Resources, Data curation, Formal analysis, Supervision, Funding acquisition, Validation, Methodology, Writing—original draft, Project administration, Writing—review and editing

**Author ORCIDs**

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Graeme J Walker [https://orcid.org/0000-0002-9392-8769](https://orcid.org/0000-0002-9392-8769)
Ethics
Animal experimentation: This study was performed in strict accordance with the recommendations of the Australian code of Practice for the care and use of animals for scientific purposes. All of the animals were handled according to approved institutional animal care and use committee of the Queensland Institute of Medical research. The protocol was approved by the Committee (A98004M). No surgery was performed. Animals were sacrificed when tumours reached 10mm in diameter, or animals were otherwise distressed.

Decision letter and Author response
Decision letter https://doi.org/10.7554/eLife.42424.018
Author response https://doi.org/10.7554/eLife.42424.019

Additional files
Supplementary files
- Supplementary file 1. Schematic representation of SNPs across the interval 4 Mb either side of Rrp15. Pink haplotypes represent the AJ/NOD allele, grey one or more of the other six founders, and clear regions were unable to be inferred. Results listed for 20 SNPs across 8 DO strains (DO1-DO11).
  DOI: https://doi.org/10.7554/eLife.42424.010
- Supplementary file 2. List of normalized gene expression values for adult mouse skin. Unprocessed raw counts based on Fragments per kilo base per million mapped reads. Table shows results for whole skin from 8 weeks old mice of the following strains; AJ, NOD, 129S, FVB, and C57BL/6
  DOI: https://doi.org/10.7554/eLife.42424.011
- Supplementary file 3. Fold change of genes at various time-points post neonatal UVR. Significantly up or down regulated genes based on analysis of either whole epidermis or whole dermis gene expression using Illumina MouseWG-6 v2.0 Expression BeadChips. At least three mice (control), and three mice (+UV) were used for each time-point.
  DOI: https://doi.org/10.7554/eLife.42424.012
- Supplementary file 4. List of significant pathways upregulated in neonatal murine skin at various times after UVR. To generate interconnected networks based on correlations, gene lists were clustered using STRING (http://string-db.org/) based on the top 500 genes differentially under or over-expressed after neonatal UVR exposure. The terms are sorted by their enrichment value shown as statistically in terms of false discovery rate.
  DOI: https://doi.org/10.7554/eLife.42424.013
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Data availability
All data generated in this manuscript are provided in the manuscript and supporting files.

The following previously published dataset was used:

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