Genome-wide association studies and genetic architecture of common human diseases

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From International Symposium on Animal Genomics for Animal Health (AGAH 2010)
Paris, France. 31 May – 2 June 2010

Abstract
Genome-wide association scans provide the first successful method to identify genetic variation contributing to risk for common complex disease. Progress in identifying genes associated with melanoma show complex relationships between genes for pigmentation and the development of melanoma. Novel risk loci account for only a small fraction of the genetic variation contributing to this and many other diseases. Large meta-analyses find additional variants, but there is current debate about the contribution of common polymorphisms, rare polymorphisms or mutations to disease risk.

Background
Genome-wide association (GWA) methods have made great progress in the last few years mapping genetic variants contributing to risk for many common complex diseases [1-3]. These methods developed from spectacular advances in genotyping technology, greater understanding of the structure of common variation in the human genome, and continued advances in computing power and software tools. Human populations have 10 to 15 million common single nucleotide polymorphisms (SNPs). Analysis of patterns of SNP variation in the human genome [4] demonstrate that a representative set of 500,000 to 1,000,000 “tagging” SNPs can sample most common variation. Current commercial SNP chips can now screen genome-wide “tagging” SNPs in a single experiment and provide an effective approach to search for genetic variants contributing to the aetiology of complex diseases.

Results and discussion
Pigmentation and melanoma
One example of mapping genes for related phenotypes is the field of pigmentation and melanoma. Melanomas ([MIM 155600]) are malignant tumours of melanocytes and susceptibility is influenced by complex relationships between genetic and environmental factors [5,6]. Risk factors include skin pigmentation (skin colour and tanning response or phototype) and the numbers of acquired melanocytic nevi or moles on the skin [5,6]. These intermediate phenotypes interact with the key environmental factor, exposure to ultraviolet light.

The incidence of malignant melanoma in Caucasian populations has increased substantially over the last 20 years [6] probably as a result of long-term changes in sunexposure.

A number of GWA studies have been conducted recently to improve our understanding of the genetics of pigmentation and contributions to melanoma risk [7-13]. Most studies have been conducted in Caucasian populations and it is important to note that variants in several pigmentation genes show significant differences in frequency, and are responsible for differences in pigmentation between ethnic groups. Therefore, care must be taken to avoid problems of population stratification in the design and interpretation of studies on pigmentation and melanoma risk.

GWA studies have identified both known and novel pigmentation genes and results have been replicated and extended by further studies including a recent meta-analysis [14]. Genes controlling pigmentation and tanning response include ASIP [MIM 600201], IRF4 [MIM 601900], KTLG [], MC1R [MIM 155555], OCA2 [MIM
Moles and melanoma risk

The most important risk factor for melanoma is the number of acquired moles [5]. Depending on the population under study, the risk for melanoma increases by 2%–4% for each additional mole counted and individuals in the top 10% of the mole count distribution have a 5- to 10- fold higher risk. Sun exposure interacts with both pigmentation and development of moles, but there is evidence for “divergent pathways” for the roles of UV radiation exposure and high mole count in development of melanoma (Whiteman 2003). Individuals with lower melanocyte proliferation and few moles develop melanoma on body sites with high cumulative UV radiation exposure while those with high melanocyte proliferation develop melanoma on body sites with lower UV radiation exposure and more moles. Total mole count has a high heritability of ~70% [18,19] and about half the variance for mole count can be attributed to a locus in the region of CDKN2A (MIM 600160) on chromosome 9 [9,18,19]. High-penetrance coding mutations in CDKN2A are reported in families with multiple melanoma cases and these families also carry greater numbers of nevi. However, the CDKN2A variants exist at population frequencies of less than 0.1%, and so explain no more than 1%–2% of melanomas in the general population. A GWA study of mole count identified common SNPs in MTAP (MIM 156540) associated both with mole count and melanoma risk [9]. MTAP is located adjacent to CDKN2A on chromosome 9p and it is not known whether the risk alleles in the 5’ region of MTAP act through direct effects on MTAP or through effects on CDKN2A. The study also demonstrated association with mole count and melanoma risk for a second locus on chromosome 22q13. The strongest association signal was for a SNP in the second intron of PLA2G6, a gene belonging to the phospholipase A2 (PLA2) super family of genes [9].

The gene IRF4 is also associated with mole count, but shows a strong gene x age interaction [20]. The T allele for rs12203592 located in intron 4 of IRF4 was associated with high mole counts and high freckling scores in adolescents, but with low mole counts and high freckling scores in adults. The C allele (associated with higher mole count in adults) was also associated with melanoma risk, most significantly with melanoma on the trunk. The gene x age interaction could easily have been missed in a single sample combining individuals of different age groups.

Effect size and missing variation

GWA studies have made good progress in identifying genes contributing to variation in pigmentation, mole development and melanoma risk. It is estimated that variants so far identified for genes influencing skin, eye and hair colour and tanning response account for about half of the melanoma risk due to pigmentation [21]. In contrast, only 2% of variation has been explained for non-pigmentation factors associated with melanoma risk including mole count. However, many of these effects are likely to act through melanocytes and CDKN2A, implicated directly in mole development and melanoma risk, has associated pigmentation effects in chickens [22]. Missense mutations in the coding region of CDKN2A are responsible for sex-linked barring, a common plumage colour characterized by black and white barred feathers. These studies illustrate the complex relationships between genes and environment in pigmentation and the development of melanoma.
risk or heritability [2,3]. One approach to this problem has been to combine results from many studies and conduct meta-analysis of results with sample sizes of over 100,000 individuals. This approach is only possible for diseases or phenotypes where many samples have been collected with the same or similar disease definitions. Some recent examples include the analysis of smoking behavior in 74,000 individuals [23], and serum lipids in >100,000 individuals [24]. These large studies have greatly increased power and each identify many novel associated variants. However, in most cases the combined results still only explain a small proportion of the genetic variation. There has been much debate about the source of the other “missing” variation. The two main possibilities are that most causal variants are not tagged well by SNPs on commercial chips (e.g. because they occur at lower frequency or are in areas of the genome for which it is difficult to develop SNP assays), or genetic contributions to disease risk are due to many variants with odds ratios so small that they do not reach formal statistical significance despite large GWA studies. Current commercial SNP chips generally target common variants. Ability to tag causal variants depends on linkage disequilibrium, in turn influenced by differences in allele frequency between markers and low frequency or rare variants will not be well “tagged” by SNP markers on many current chips.

Contribution of rare variants

Contribution to disease risk in the population is a function of allele frequency and also of effect size for the risk allele. Rare disease associated variants not tagged by current chips can only be the source of missing heritability if the risk alleles have large effects. Re-sequencing of genomic regions uncovers new variation and there are a number of examples where rare variants contribute to risk for common traits. Rare variants in CDKN2A discussed above contribute to melanoma in high risk families, but explain little of the population prevalence for this disease. GDF9 is expressed in human oocytes and plays important roles in growth and selection of ovarian follicles. A search for GDF9 variants in mothers of spontaneous DZ twins identified three novel deletions and four mis-sense alterations [25,26]. Taken together, the frequency GDF9 variants were significantly higher in mothers of DZ twins compared with controls [25,26]. However, the frequency of the variants is low (less than 4% for all variants) and the contribution of these variants to the overall incidence of twinning is small. Re-sequencing a candidate gene for type 1 diabetes detected new variants at ~1% frequency that in total contributed more to variation in risk in the population than a single common variant in the same gene detected by a previous GWA study [27]. Recently, GWA identified common variants APOA5, GCKR, LPL and APOB associated with hypertriglyceridemia (HTG, [28]). Resequencing of these genes revealed a significantly higher burden of rare missense or nonsense variants in individuals with HTG, compared to controls corresponding to a carrier frequency of 28.1% of affected individuals and 15.3% of controls. Common genetic variants in seven HTG-associated loci explained ~20% of total variation in HTG diagnosis, and the rare genetic variants in four HTG-associated loci explained ~1% of variation. Therefore, both rare and common variants in the same genes can influence disease risk. Based on current examples, the contributions from rare or low frequency variants are similar to common variants and much variation in genetic contributions to disease risk is still “missing”.

Common variants of small effect explain missing heritability

Most GWA studies have examined evidence for association SNP by SNP. An alternative approach is to analyse data for all SNPs together to estimate the proportion of trait variance accounted for by all common variation “tagged” by the SNPs on current commercial SNP chips. This is possible because the distant genetic relatedness of individuals can be estimated from dense SNP data. Once the degree of relatedness is established, it can be compared to phenotypic similarity between the individuals. This method was developed and used to estimate the genetic contribution to variation in height independent of the usual assumptions required to estimate heritability using family data [29]. Using this approach, the percentage of phenotypic variation explained by common SNPs was 45%.

This is less than the 80% of phenotypic variance due to additive genetic effects based on the estimated heritability. However, the SNPs sampled on the arrays may not be in complete LD with the causal variants and this might influence the results. If the estimate is corrected first for the sampling error from using a finite number of SNPs with genotype data, the corrected estimate for variance explained by causal variants is 54% (assuming the same structure of linkage disequilibrium between causal variants and common SNPs sampled on the arrays). In addition, if the causal variants tend to have lower minor allele frequencies than SNPs on the arrays, we would expect lower LD between genotyped SNPs and causal variants [29]. When this is taken into account, the estimated contribution of phenotypic variation explained was 84%. The standard error for this estimate is large and it does not prove that causal variants do have lower allele frequencies than tagging SNPs used on the chips. However, if this were the case, most of the phenotypic variation in height due to additive genetic effects could be explained by many common variants.
with small effects [29]. Whether this applies only to human height or more generally remains to be seen.

Conclusions

GWA studies have identified a large number of variants associated with a range of human traits and common diseases. However, the sizes of effects on disease risk are typically small. Combining results across many studies increases the power to detect risk variants and sequencing is uncovering rare variants with modest contributions to a number of diseases. The emerging view from all these studies is a spectrum of many variants with small effects explaining genetic contributions to disease risk.

The discoveries provide new insights into the biology of many diseases with a number of variants located in genes that contribute to biological pathways not previously considered to be involved in disease, or located in regions that do not contain known protein-coding genes. Some examples like the effects of IRF4 on mole count show interactions that would reduce estimated effects size from large combined studies. Therefore one important outcome of GWA studies will be to use knowledge gained to evaluate genetic contributions to disease sub-classes, disease heterogeneity and co-morbidity for different diseases. The next challenge is how to translate these discoveries into better diagnostic practices, prevention and treatments.

Acknowledgements

This article has been published as part of BMC Proceedings Volume 5 Supplement 4, 2011: Proceedings of the International Symposium on Animal Genomics for Animal Health (AGAH 2010). The full contents of the supplement are available online at http://www.biomedcentral.com/1753-6561/5?issue=S4.

Competing interests

The authors declare that they have no competing interests.

Published: 3 June 2011

References

1. Hindorff LA, Sethupathy P, Junkins HA, Ramos EM, Mehta JP, Collins FS, Manolio TA: Potential etiologic and functional implications of genome-wide association loci for human diseases and traits. Proc Natl Acad Sci U S A 2009, 106(23):9362-9367.
2. McCarthy MI, Abecasis GR, Cardon LR, Goldstein DB, Little J, Ioannidis JP, McShane LM: Genome-wide association studies for complex traits: consensus, uncertainty and challenges. Nature Reviews Genetics 2008, 9(5):356-369.
3. Visscher PM, Montgomery GW: Genome-wide association studies and human disease: from trickle to flood. JAMA 2009, 302(18):2028-2029.
4. Frazer KA, Ballinger DG, Cox DR, Hinds DA, Stuve LL,GBSD RA, Belmont JW, Boudreau A, Hardenbol P, Leal SM, et al: A second generation human haplotype map of over 3.1 million SNPs. Nature 2007, 449(7164):851-861.
5. Bauer J, Garbe C: Acquired melanocytic nevi as risk factor for melanoma development. A comprehensive review of epidemiological data. Pigment Cell Res. 2003, 16(3):297-306.
6. Meyle KD, Guildberg P: Genetic risk factors for melanoma. Human Genetics 2009, 126(4):499-510.
7. Bishop DT, Demenais F, Ilies MM, Harland M, Taylor JC, Corda E, Randerson-Moor J, Arkin JF, Avril MF, Azali E, et al: Genome-wide association study identifies three loci associated with melanoma risk. Nat Genet 2009, 41(8):920-925.
8. Brown KM, Nacagoe S, Montgomery GW, Craig DW, Zhao ZZ, Inadomi K, Henders AK, N H, Campbell MJ, Stark MS, et al: Common sequence variants on 20q11.22 confer melanoma susceptibility. Nat Genet 2008, 40:838-840.
9. Faltchi M, Bataille V, Hayward NK, Duffy DL, Bishop JA, Pastinen T, Cervino A, Zhao ZZ, Deloukas P, Saranbo N, et al: Genome-wide association study identifies variants at 9p21 and 22q13 associated with development of cutaneous nevi. Nat Genet 2009, 41(8):915-919.
10. Gudbjartsson DF, Sulem P, Stacey SN, Goldstein AM, Rafnar T, Sigurjonsdottir B, Benediktsson KR, Thorisdottir K, Ragnarsson R, Sveinsdottir SG, et al: ASIP and TYR pigmentation variants associate with cutaneous melanoma and basal cell carcinoma. Nat Genet 2008, 40:886-891.
11. Han J, KRAFT P, Nan H, Guo Q, Chen C, Qureshi A, Hankinson SE, Hu FB, Duffy DL, Zhao ZZ, et al: A genome-wide association study identifies novel alleles associated with hair color and skin pigmentation. PLoS Genet 2008, 4(5):e1000074.
12. Sulem P, Gudbjartsson DF, Stacey SN, Helgason A, Rafnar T, Jakobsson M, Steinberg S, Gudjonsson SA, Palsson A, Thorleifsson G, et al: Two newly identified genetic determinants of pigmentation in Europeans. Nat Genet 2008, 40:835-837.
13. Sulem P, Gudbjartsson DF, Stacey SN, Helgason A, Rafnar T, Magnusson KP, Manolescu A, Karason A, Palsson A, Thorleifsson G, et al: Genetic determinants of hair, eye and skin pigmentation in Europeans. Nat Genet 2007, 39(12):1443-1452.
14. Genetebhif MR, SH N, Landt M: Genome-wide association studies of pigmentation and skin cancer: a review and meta-analysis. Pigment Cell Melanoma Research 2010, Epub 14 June 2010.
15. Duffy DL, Box NF, Chen W, Palmer JS, Montgomery GW, James MR, Hayward NK, Martin NG, Sturm RA: Interactive effects of MCHR1 and OCA2 on melanoma risk phenotypes. Human Molecular Genetics 2004, 13:447-451.
16. Elberg H, Troelsen N, Nielsen M, Mikkelsen A, Mengel-Froim J, Kjaer KW, Hansen L: Blue eye color in humans may be caused by a perfectly associated founder mutation in a regulatory element located within the HERC2 gene inhibiting OCA2 expression. Hum Genet 2008, 123(2):177-187.
17. Sturm RA, Duffy DL, Zhao ZZ, Leite FP, Stark MS, Hayward NK, Martin NG, Montgomery GW: A single SNP in an evolutionary conserved region within intron 86 of the HERC2 gene determines human blue-brown eye color. Am J Hum Genet 2008, 82(2):424-431.
18. Falchi M, Spector TD, Perks U, Kato BS, Bataille V: Genome-wide search for nevus density shows linkage to two melanoma loci on chromosome 9 and identifies a new QTL on 5q31 in an adult twin cohort. Human Molecular Genetics 2006, 15(20):2969-2979.
19. Zhu G, Montgomery GW, James MR, Trent JM, Hayward NK, Martin NG, Duffy DL: A genome-wide scan for nevus count: linkage to CDKN2A and to other chromosome regions. Eur J Hum Genet 2007, 15(4):104-102.
20. Duffy DL, Iles MM, Glass D, Zhu G, Barrett J, Hixon V, Zhao ZZ, Sturm RA, Saranbo N, Hammond C, et al IRF4 variants have age-specific effects on nevus count and predispose to melanoma. Am J Hum Genet 2010, 87:6-16.
21. Duffy DL, Zhao ZZ, Sturm RA, Hayward NK, Martin NG, Montgomery GW: Multiple Pigmentation Gene Polymorphisms Account for a Substantial Proportion of Risk of Cutaneous Malignant Melanoma. Journal of Investigative Dermatology 2010, 130:520-528.
22. Hellstrom AR, Sundstrom E, Gunnarsson U, BedHorn B, Tavier-Bochard M, Honaker CF, Sahliqvist AS, Jensen P, Karpas V, Siegel PB, et al: Sex-linked barring in chickens is controlled by the CDKN2A /B tumour suppressor locus. Pigment Cell Melanoma Research 2010, 23(4):521-530.
23. Thorgeirsson TE, Gudbjartsson DF, Surakka I, Vink JM, Amin N, Geller F, Sulem P, Rafnar T, Esco T, Walter S, et al Sequence variants at CHRNB3, CHRNAB and CYP2A6 affect smoking behavior and the risk of lung cancer. Nat Genet 2010, 42:448-453.
24. Teslovich TM, Musunuru K, Smith AV, Edmondson AC, Stylianou IM, Kosik M, Pirruccello JP, Ripatti S, Chrom sub ctal J, et al: Biological, clinical and population relevance of 95 loci for blood lipids. Nature 2010, 466:707-713.
25. Montgomery GW, Zhao ZZ, Marsh AJ, Mayne R, Treloar SA, James MR, Martin NG, Boomsma DI, Duffy DL: A deletion mutation in GDF9 in sisters with spontaneous DZ twins. Twin Research 2004, 7:548-555.

26. Palmer JS, Zhao ZZ, Hoekstra C, Hayward NK, Webb PM, Whiteman DC, Martin NG, Boomsma DI, Duffy DL, Montgomery GW: Novel variants in growth differentiation factor 9 in mothers of dizygotic twins. Journal of Clinical Endocrinology and Metabolism 2006, 91:4713-4716.

27. Nejentsev S, Walker N, Riches D, Egholm M, Todd JA: Rare variants of IFIH1, a gene implicated in antiviral responses, protect against type 1 diabetes. Science 2009, 324(5925):387-389.

28. Johansen CT, Wang J, Lanktree MB, Cao H, McIntyre AD, Ban MR, Martins RA, Kennedy BA, Hassell RG, Visser ME, et al: Excess of rare variants in genes identified by genome-wide association study of hypertriglyceridemia. Nat Genet 2010, 42(8):684-687.

29. Yang J, Benyamin B, McEvoy BP, Gordon S, Henders AK, Nyholt DR, Madden PF, Heath AC, Martin NG, Montgomery GW, et al: Missing heritability of human height explained by many genes of small effect. Nat Genet 2010, 42(8):655-659.

Cite this article as: Montgomery: Genome-wide association studies and genetic architecture of common human diseases. BMC Proceedings 2011 5(Suppl 4):S16.

doi:10.1186/1753-6561-5-S4-S16

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