Molecular Genetics of Alcohol Dependence and Related Endophenotypes

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Abstract: Alcohol dependence is a worldwide public health problem, and involves both environmental and genetic vulnerability factors. The heritability of alcohol dependence is rather high, ranging between 50% and 60%, although alcohol dependence is a polygenic, complex disorder. Genome-wide scans on large cohorts of multiplex families, including the collaborative study on genetics of alcoholism (COGA), emphasized the role of many chromosome regions and some candidate genes. The genes encoding the alcohol-metabolizing enzymes, or those involved in brain reward pathways, have been involved. Since dopamine is the main neurotransmitter in the reward circuit, genes involved in the dopaminergic pathway represent candidates of interest. Furthermore, gamma-amino-butyric acid (GABA) neurotransmitter mediates the acute actions of alcohol and is involved in withdrawal symptomatology. Numerous studies showed an association between variants within GABA receptors genes and the risk of alcohol dependence.

In accordance with the complexity of the “alcohol dependence” phenotype, another field of research, related to the concept of endophenotypes, received more recent attention. The role of vulnerability genes in alcohol dependence is therefore re-assessed focusing on different phenotypes and endophenotypes. The latter include brain oscillations, EEG alpha and beta variants and alpha power, and amplitude of P300 amplitude elicited from a visual oddball task.

Recent enhancement on global characterizations of the genome by high-throughput approach for genotyping of polymorphisms and studies of transcriptomics and proteomics in alcohol dependence is also reviewed.

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INTRODUCTION

Alcohol dependence is a complex addictive disorder, affecting 5.4% of the general population during lifetime [1]. Alcohol dependence (Online Mendelian Inheritance in Man #103780) represents a complex and heterogeneous phenotype, with behavioural, as well as psychological, pharmacological, and medical components. Aggregation and adoption researches support the role of a genetic component in alcohol dependence. Twin studies estimate that the additive heritability ranges between 50% and 60% [2]. However, the exact transmission mode remains unknown, but more than one gene is likely to be involved (i.e. pauci- or poly-genic disorder).

This paper focuses on current studies on the genetics of alcohol dependence. After an overview of genetic approaches, including genome-wide scans and candidate genes analyses, we will review post gene methods, such as expression profile studies of genes differentially expressed in phenotypes relevant for alcohol dependence.We will particularly discuss the results from two emerging fields, namely genomic and proteomic approaches of alcohol dependence.

GENETICS OF ALCOHOL DEPENDENCE

The identification of genes involved in alcohol initiation, use, abuse and dependence has been investigated by two complementary genetic methods, namely (i) genome wide scan analyses, with no a priori assumptions about which genes might be involved, and (ii) the genotyping of candidate genes of interest. About fourteen-hundreds papers came out from the PubMed/NCBI database with the key words “alcohol, dependence, gene”. This review is not an exhaustive view of linkage studies on alcohol dependence, neither a cover of all genes possibly affecting the risk for alcohol dependence. We have rather decided to focus on particularly relevant linkage and candidate genes studies. For a more complete review refers to Goldman et al. [3] and Ramoz et al. [4]. We also excluded gene environment interactions of our review, and refer the reader to Gorwood et al. [5].

Genome-Wide Scans of Alcohol-Dependence

Recently, linkage studies have identified several chromosomal regions, supporting the involvement of loci on chromosomes 1, 2, 4, 5, 6, 7, 12, 14, 16 and 17 in alcohol dependence [3, 4, 6]. Most of these results were found on alcohol dependent families from the collaborative study on the genetics of alcoholism (COGA). The COGA consists of large pedigrees ascertained using probands who met diagnostic not only for alcohol dependence from the Diagnostic
Candidate Genes

Candidate Genes in the Metabolism of Alcohol

ADH Genes

The chromosome 4q region surrounding the alcohol dehydrogenase (ADH) gene was linked in a genome-wide scan with the maximum numbers of drinks ever consumed in 24 hours, a phenotype associated with alcohol dependence [28]. The role of ADH is to convert alcohol into acetaldehyde. The seven ADH genes cluster on chromosome 4q (from 5′ to 3′ region: ADH7-ADH1C-ADH1B-ADH1A-ADH6-ADH4-ADH5). However, only two exhibit functional polymorphisms, ADH1B*2 and ADH1C*2 alleles, and confer an increase in alcohol degradation rate [29, 30]. The class I enzymes (encoded by ADH1A, ADH1B and ADH1C) and the class II enzyme (encoded by ADH4) are significant contributors to ethanol metabolism, at least in vitro [30]. ADH1B*2 is relatively common among Asian population [31], as well as in Israeli Jews [32]. It is associated with a diminution in alcohol consumption and therefore has a protective effect against alcohol dependence [31]. ADH1C*2 has protective effects on liver cirrhosis and alcohol chronic pancreatitis [33], an is also associated with a lower rate in alcohol dependence among Asian population, maybe because ADH1B and ADH1C are in linkage disequilibrium[34]. Indeed, ADH1B and ADH1C are separated by a distance of only 15kbp in the cluster of ADH genes. Contrasting with their major effect in vitro, ADH1C and ADH1B genes have minor effects in vivo, suggesting a possible role of other genes of the ADH cluster. Indeed, the effect of the quantitative trait loci (QTL) linked to the ADH region gene account for 64% of the genetic variance in alcohol metabolism [35]. For example, ADH4 contribute to 30% of the total ethanol oxidizing capacity of the liver [30]. Recent findings suggest evidences of association with alcoholism [30, 36], as well as with personality traits [37], although among non-coding SNPs. Similarly, a haplotype of the ADH7 gene contribute in the inter-individual variation in alcohol metabolism [38], and is associated with alcohol dependence [39], drug dependence [40] as well as with personality traits [41].

ALDH Genes

Acetaldehyde dehydrogenase (ALDH) genes encode a protein that acts as a tetramer to oxidize acetaldehyde [3,4]. The blockage of ALDH with disulfiram leads to the accumulation of acetaldehyde in the blood, and results in a flushing syndrome, that is a reddening of the face, sudden sweating, vomiting and headache. This syndrome can also occur in subjects carriers of the ALDH2*2 variant, coding for an inactive enzyme. The deficient variant allele acts dominantly, thus ALDH2*1/ALDH2*2 heterozygous carriers have reduced enzyme activity, but the flushing reaction is more severe and immediate in ALDH2*2 homozygous carriers [42]. The ALDH2*2 is associated with a reduced rate of alcohol dependence in Asian population [43], and in both European and African Americans [44].

Interestingly, ADH1B and ALDH2 genes have an additive effect on the risk of flushing, with no epistatic interaction involved, that is the protective effects conferred by one gene are independent of those conferred by the other [44].

Candidate Gene Involved in the Rewarding Circuits

Ethanol modulates several neurotransmitter systems involved in alcohol initiation, tolerance, preference, consumption, abuse and dependence [4]. These relationships interplay with neurocognitive functions, personality traits related either to alcohol seeking process or to subjective effects of...
ethanol. Regarding the neurobiological pathways involved in these processes, they are linked to the reward pathway, which mainly consisted of the dopaminergic projection of the ventral tegmental area on the nucleus accumbens. This dopaminergic pathway is modulated by several other neurotransmitters, including the GABA.

**Dopamine Pathway**

The ability of alcohol to increase brain dopamine concentration in the mesolimbic pathway has a key role in its reinforcing effects. Dopamine is also involved in withdrawal physiopathology, as suggested by the lower levels of striatal D2 receptors during alcohol withdrawal [45]. Several genes are involved in the neurobiological dopaminergic pathway, which have led researchers to assess different candidate gene polymorphisms. For instance, the single nucleotide variant TaqIA rs1800497 of the dopamine D2 receptor. **DRD2** gene, has been considered as a vulnerability gene for alcoholism in more than 40 studies, but with conflicting results, as suggested by meta-analysis [46, 47]. Interestingly, and in accordance with the role of dopamine in the detection of potential reward facets of external stimuli, the A1 allele carriers of the rs1800497 learned to avoid actions with negative consequences less efficiently [48]. Therefore, they are less likely to learn from the negative consequences of their alcohol consumption, and thus may be more likely to become dependent. This indirect involvement could partly explain the discrepant results observed in previous association studies. On the other hand, recent dense mapping of the DRD2 locus suggests association with adjacent genes, namely ANKK1, TTC2 and NRCAM1 [49, 50]. In fact, TaqIA variants change the amino acid in the ANKK1/X-kinase gene [51], and might confer functional biological consequences in alcohol dependence.

Recently, association between alcohol dependence and a SNP in the **DRD1** gene has been shown in a French population [52]. This is convergent with a previous association between DRD1 (among 5 genes) in a Korean population [53]. While there is evidence for the role of D3 receptor in alcohol intake in animal [54], genetic studies of dopamine receptor D3 and D4 genes failed to show associations with vulnerability to alcoholism [55-57].

The **DAT1** gene encodes the dopamine transporter (DAT) and represents a promising candidate gene in the dopaminergic system. This transporter is mainly located at the synapse of the presynaptic neuron and is responsible for dopamine reuptake. Human genetic studies report associations between variants of the **DAT1** gene and alcohol dependence with both positive [58, 59] and negative [60, 61] results. Indeed, the A9 variant of the **DAT1** is associated with more severe withdrawal symptomatology [62], and complications [63]. More specifically, this allele is associated with withdrawal seizures, probably through the modulation of neuronal excitability [64].

**Gamma-Amino-Butyric Acid (GABA) Pathway**

GABA is the major inhibitory neurotransmitter in the brain. The GABA receptor is a pentameric protein containing two α, two β, and one γ or δ subunit among 19 isoforms. This arrangement allows considerable diversity of the GABA receptor, but their observed number is in fact limited. Alcohol is an agonist of the GABA_A receptor, particularly receptors containing the α4 or α6 subunits in combination with β2 or 3 subunits and the δ subunit [65]. The GABAergic activity is reduced during alcohol withdrawal and could account for a part of the symptomatology. For example, in a mouse model, a mutant of the γ2 subunit is associated with an increased severity of ethanol withdrawal [66].

**GABRA6** variants were associated with alcohol dependence in different populations, including a Scottish [67], a German [68], a Finn and a Native American sample [66]. Variations in **GABRA2** were associated with alcohol dependence [69] as well as with illicit drug dependence [70] in the COGA sample, and this association was recently replicated in an independent sample [71].

**HUMAN POST-GENE STUDIES AND ANIMAL MODELS: SEARCH FOR ALCOHOL DEPENDENCE CANDIDATE GENES**

Since almost a decade now, the traditional tools used in human genetic studies to identify alcohol dependence genes, genome-wide scans and candidate gene associations, have been helped by the high-throughput characterization of the expression patterns of the genes (transcriptome studies) and the proteins (proteome studies) from tissue samples. This tissue can be extracted from specific brain regions, of patients compared to controls, as well as from animal used as models of alcohol dependence, from the yeast to the murine models. Indeed, although there are several studies describing global gene expression changes associated with alcohol dependence, changes in the expression levels of proteins has not been characterized until recently. Characterization of the liver, brain and serum proteome is the next step toward the investigation of alcohol-dependent patients to discover new biological markers as well as prognostic tools.

**Transcriptome Studies**

Whole-brain gene expression databases of five genetic mouse models with different alcohol consumption were examined [72]. A meta-analytic approach across data sets allowed the identification of 36 candidate genes for high and low amounts of alcohol consumption. Some of these genes were located within quantitative trait loci (QTLs) for human alcohol dependence susceptibility. Among them, 11 genes were clustered in only three gene families: GSTM (glutathione S-transferase activity), S100 class calcium-binding, and cytokine/proinflammatory activity (IL-1). Using microarray analyses of cerebellar tissue from mutant and wild-type mice exposed to alcohol, 109 genes were associated with either the development or the resistance to tolerance, or with the effects of chronic alcohol administration [73]. Specifically, six genes were both relevant to ethanol response and had modifications in micro array expression profiles further confirmed by quantitative real-time PCR. Of them, Twik-1, coding for a potassium channel, was associated with tolerance, and JunB and Nur77, members of an immediate early gene family and of the nuclear receptor family respectively, had an increased expression that persisted a long time after alcohol ingestion [73]. These two recent works illustrated the diversity of many gene expression studies on central nervous system from in vitro cell lines and in vivo animal models,
especially in mice and rats [74-77]. It is important to note that transcriptome studies are usually coupled with phylogenetic and functional biological pathway studies for the analysis of candidate genes, using GENE ONTOLOGY and IN-GENUITY packages, and may also integrate human genetic data in a more global approach called functional genomics, as illustrated in [74, 76, 77].

Due to the difficulty to have access to human post-mortem brain samples, the number of differential display and transcriptome studies on alcohol dependence, as well as the sample size of their studies, are modest. Nevertheless, several works have been published in the last years on independent samples of different brain regions, including temporal cortex, frontal cortex and nucleus accumbens of alcohol dependent patients and controls [78-82]. One of the first microarray gene expression studies identified a novel neuron-specific gene, hNP22(TAGLN3), over-expressed in the superior cortex of patients [78]. Significant differences in the expression levels were observed between cases and controls for genes involved in mitochondrial, ubiquitination and signalling pathways in the temporal cortex [79]. Genes encoding mitochondrial and metabolism proteins, as well as genes associated with neuroprotection and apoptosis, were also found in the prefrontal cortex of cases and not at the same level in controls [80]. Nucleus accumbens study revealed genes involved in the vesicle formation, in the structure, adhesion and migration of cells in the same patients [80]. Another expression analysis of other samples from the frontal cortex also showed dysregulation of genes implicated in metabolism, energy production, cell survival and communication [81]. This study was extended to additional cases and controls, and found altered expression of genes involved in myelination, ubiquitination, apoptosis, cell adhesion and neurogenesis [82].

The different pathways identified by the modifications of gene expression should help in the understanding of the molecular mechanisms of alcohol dependence, in particular by allowing the characterisation of protein levels to reveal new biological markers, and therefore increase the chance to further understand the pathophysiology of this psychiatric disorder.

**Proteome Studies**

Different methods in proteomics, such as liquid chromatography, mass spectrometry (MALDI-TOF and SELDI-TOF), and two-dimensional gel electrophoresis, have been recently carried out to identify and quantify proteins in the field of addiction. They have been used on cell lines, animal models, as well as on post-mortem tissues, in particular brain, of patients to characterise changes in protein expression in response to alcohol consumption, as presented in recent reviews [4, 83, 84]. Three recent works in this field could be emphasized, two on animal models and one in human post-mortem brain samples. In the first study, proteomic approach has been performed on liver from rats of inbred alcohol-prefering line (iP) exposed to alcohol or not [85]. The authors found 113 proteins with changes of expression levels. Seventy-one percent were enzymes and most of them are involved in metabolism pathways, including glycolysis, gluconeogenesis or fatty acid oxidation. These results, in particular the up-regulation of enzymes involved in antioxidant activity, are highly consistent with the known effects of alcohol on the liver. In the second study using mass spectrometry and confirmation by immunoblot analysis, levels of proteins in the serum of primates (cynomolgus monkey), trained to self-administer ethanol, were compared between “drinking” and “abstinent” for periods of 6 months [86]. They observed a significant increase in apolipoproteins A1 and AII. This observation is concordant with the detection of apolipoprotein in the serum of human ethanol consumption [87], and shows that this primate model represents a remarkable model to identify biological markers and maybe helpful to characterize the pathophysiology of alcohol dependence. In the third work, protein expression profiles from the white matter of the BA9 region were examined between alcohol dependent patients and controls [88]. Most of the differences were observed when comparing controls to severe alcoholic patients with complications (hepatic cirrhosis). Furthermore, some but not all spots of detected protein returned to control level in recovery alcohol dependent patients. This is of interest because these proteins could be involved in the regeneration of white matter after a prolonged abstinence. A total of 60 proteins were dysregulated in patients with alcohol dependence. Among these proteins, the level of hNP22 was significantly decreased (~3-fold) in complicated patients [88]. This biomarker was previously identified by its gene expression level in the superior cortex of patients [78] and corresponds to a hydrophilic, cytoplasmic protein involved in the cytoskeleton of the cell. Others proteins playing a role in the metabolism, mitochondria, energy production and ubiquitination were characterized [88], and were also previously identified by gene expression analysis.

Thus, a key area for proteomics consists in biomarker discovery that is the identification of protein expression which correlates with disease, treatment outcome or global prognosis, thus allowing patient-tailored therapy. In that view, the results presented above showed promising results. Furthermore, some biological markers were also found by gene profiling, and vice versa, which reinforce their probable implication in alcohol dependence. Finally, the last presented study point out the fact that the clinical features and phenotype of the patients need to be the most well documented to perform studies on comparable patients and homogenous groups. This lead to the novel strategy followed in genetic studies looking for susceptibility to psychiatric disorder which consists to identify specific traits that help to increase the homogeneity of the group of patients, when the disorder has a broad spectrum phenotype, such as alcohol dependence.

**ENDOPHENOTYPIC APPROACH**

An endophenotype is a biological or a psychological characteristic transmitted with a disease, with a strong biological substrate and a higher heritability than the disease itself [89]. The required qualities of an endophenotype also include sensitivity and specificity, heritability, presence in unaffected relatives, state-independence, biological plausibility, sound psychometric properties and feasibility. The electrophysiological related features constitute the most studied endophenotype in alcohol dependence, and associates
electroencephalography (EEG) variation and event-related potentials (ERPs).

Brain Oscillations, EEG Alpha and Beta Variants and Alpha Power

EEG features are partially heritable and relatively stable among lifespan [90]. The alpha rhythm (7.5-12 Hz) is the posterior wave form, detected during alert relaxation. One common variant in EEG pattern concerns the alpha waves, since low voltage alpha (LVA) have previously shown a pattern of inheritance compatible with an autosomal dominant mode of inheritance [91]. LVA is associated with a familial history of alcohol dependence in subjects with no personal history of alcohol related problems. This finding was further confirmed on several ethnicities [92]. The beta waves (13–28 Hz) are also significantly different between alcohol dependent patients and controls. Moreover, the offspring of alcohol-dependent males also exhibited this variation [93], and was therefore used as an endophenotype in several genetic studies, in linkage analysis [94,95] and in candidate gene association [16, 20, 69].

Amplitude of P300 Elicited from a Visual Oddball Task

The P300 amplitude reflects the allocation of attention, with the larger amplitude, the more attention allocated. It is measured during an oddball paradigm where patients are asked to detect target stimuli, and to ignore standard and non-target rare events. It is a complex wave, with at least two distinct components or ERPs, namely the P3a and the P3b waves. The P3a, the early component, is elicited by novel and unexpected stimuli, whereas the later P3b wave is elicited when context updating is necessary.

The P300 is a highlyheritable phenotype, since the estimatedheritability ranges between 0.30 and 0.81 depending on the studies [96], and was therefore used as an endophenotype also in alcohol related disorders. Indeed, the P300 amplitude of the ERPs is reduced in the sons of alcoholic fathers, with no previous exposure to alcohol [97]. This variation was observed in several desinhibitory disorders, such as conduct disorder, attention-deficit hyperactivity disorder, oppositional defiant disorder, and antisocial personality disorder [98], suggesting that the P300 is associated with an intermediate phenotype, such as impulsivity, rather than with alcohol dependence itself.

CONCLUSIONS

Our current knowledge on the neurobiological substrate of alcohol dependence leads to propose a high number of candidate genes. Association and linkage studies of the genes encoding proteins involved in alcohol metabolism (e.g. ADH, ALDH) and in the rewarding circuits, including dopamine (DRD2, DAT1) and GABA (GABRA2, GABRA6) neurotransmitter biological pathways, are the most replicated findings. It is important to note that novel candidate genes for alcohol dependence were identified from animal models. Thus, the drosophila hangover gene was found as required for normal development of ethanol tolerance in the fruit flies [99]. Human HANGOVER/ZNF699 gene polymorphisms were associated in a large population of Irish alcohol dependent patients compared to matched controls [100].

Genetic studies in alcohol dependence also recently took advantage of novel technology for high-throughput genotyping based on microarrays. Thus, association genome scanning, based on the simultaneous genotyping of over 100,000 SNPs, was carried out on the COGA [101]. Convergent loci and associations, as previously reported, were found. Furthermore, novel loci were identified confirming the high definition of this new tool in screening capacity [101]. Recently, rare structural variants, named copy number variants (CNVs), which correspond to micro-duplications or micro-deletions of 100 kb at least leading to duplicate or disrupt genes, were identified by comparative genomic hybridization in several psychiatric disorders [102, 103]. Thus, search for CNVs should be one of the next future directions in the molecular genetic of alcohol dependence.

Genomic functional, including gene and protein expression profiling studies, on cell lines, animal models and human post-mortem samples confirmed that several biological pathways are involved in the vulnerability to alcohol preference, and opened avenues for patient-tailored treatment, in both pharmacogenomic and clinical proteomic. This genomic functional approach has to be combined with genetic analyses, and have to be performed on large number of patients, assessing a large set of clinical features and phenotypic information, also including endophenotypes to narrow the associated gene, or biomarker, to a specific step in the pathway to alcohol dependence. Finally, pharmacogenomics is the next issue in the comprehension of the molecular biology of alcohol dependence.

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