Transcriptome profiling of whisker follicles in methamphetamine self-administered rats

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Methamphetamine (MA) is a highly addictive psychostimulant that disturbs the central nervous system; therefore, diagnosis of MA addiction is important in clinical and forensic toxicology. In this study, a MA self-administration rat model was used to illustrate the gene expression profiling of the rewarding effect caused by MA. RNA-sequencing was performed to examine changes in gene expression in rat whisker follicles collected before self-administration, after MA self-administration, and after withdrawal sessions. We identified six distinct groups of genes, with statistically significant expression patterns. By constructing the functional association network of these genes and performing the subsequent topological analysis, we identified 43 genes, which have the potential to regulate MA reward and addiction. The gene pathways were then analysed using the Reactome and Knowledgebase for Addiction-Related Gene database, and it was found that genes and pathways associated with Alzheimer's disease and the heparan sulfate biosynthesis were enriched in MA self-administration rats. The findings suggest that changes of the genes identified in rat whisker follicles may be useful indicators of the rewarding effect of MA. Further studies are needed to provide a comprehensive understanding of MA addiction.

Addiction is a representative disorder of the brain’s reward system and occurs over time from high chronic exposure to an addictive stimulus, such as an abuse drug. The defining feature of addiction is compulsive and out-of-control drug use. The uncontrolled drug use may lead to criminal or anti-social behaviour, violence, physical dependence, or psychological addiction.

Methamphetamine (MA), the second most abused illegal psychostimulant, affects the central nervous system. MA has pervasive effects not only on the dopaminergic system, but also on diverse neurotransmitter systems, such as noradrenergic and serotonergic systems, in the brain. Converging evidence suggests that chronic MA use is associated with immunological dysfunction and impaired neurogenesis in the limbic dopaminergic system. The reported neurotoxicity induced by MA might be related to reactive oxygen species, apoptotic processes, and excitotoxicity, which might be a major cause of neurodegenerative disorders, in the dopaminergic and serotonergic systems. Clinical observation suggests that MA causes long-lasting injury to the brain. Moreover, diverse clinical symptoms, such as depression, suicidal ideation, and psychotic behaviour, are observed in people who are addicted to MA.

MA abuse and addiction have become a serious worldwide issue since the 1990s. Ease of manufacture and distribution is the major factor linked to increased use of MA. In Republic of Korea, MA is the most widely used illegal drug, and 100% of persons treated for drug problems in the country are MA abusers. In particular, MA abuse is a growing serious problem in youths; thus, investigation on MA abuse is needed to understand the seamy side of MA. Therefore, testing for MA abuse is crucial for assessing the reality of intoxication and evaluating the level of drug impairment.

Testing for MA abuse is largely dependent on biofluids, such as urine, blood, and plasma. Urine is the most used traditional specimen; urine testing is a valuable measure of drug use in patients who use the drug regularly. However, urine testing has limitations for those with moderate drug use because of the short drug detection time. In contrast, the use of the hair has several advantages, such as ease of collection, noninvasiveness, and a

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long detection window (months to years), for toxicological analysis. Interestingly, a recent report suggests that scalp hair follicles can be used to find genes associated with brain diseases because the brain is an ectodermal tissue and shares the developmental origins with scalp hair follicle. Moreover, the hair follicle system, which is a well-coordinated miniorgan within the skin, has been used to detect stressors, such as neurohormone, neurotransmitters, and cytokines, as well as to diagnose traumatic brain injury. Therefore, it could be valuable to study changes in gene expression in hair follicles, instead of brain tissues, in order to identify novel indicators related to the rewarding effect of MA in living donors.

The development of addiction is accompanied by changes in the signal transduction pathway and cellular gene expression in the brain. Many addiction-related genes and pathways have been identified, however, the exact mechanism underlying MA addiction has not been fully elucidated. Previous studies have implied that exposure to MA causes complex molecular alterations; therefore, a comprehensive multigene level approach is needed to understand the drug addiction mechanisms, and it is important to identify the principal molecular network underlying MA addiction by analysing alteration in gene networks.

RNA-sequencing (RNA-seq) is transcriptome profiling and uses deep-sequencing technologies. RNA-seq is used to investigate the expression of unknown and known candidate genes with higher sensitivity than microarray. Genes that are significantly upregulated or downregulated can be determined in certain conditions by quantifying changes in RNA expression between different conditions. However, RNA-seq alone may not provide pragmatic results in a multigenic disease, such as drug addiction, because of extensive raw data; therefore, data management and computational screening are needed. The integration of bioinformatics and biological network analysis enables RNA-seq to define specific groups of genes that are related to drug addiction behaviour. Recent advances in bioinformatics facilitate analysis of complex signalling transduction in cells through investigation of multiple types of interaction networks. Biological network analysis is a powerful tool using computable networks in the biological system. Gene-gene association network analysis provides a novel method of analyzing a large number of differentially expressed genes by classifying them into different gene groups based on their function or signalling pathway annotation. In the network analysis, degree, which is the number of edges connecting a node, and betweenness centrality (BC), which is how central the node is to the network, are used to show the importance of the genes.

In this study, RNA-seq was used to illustrate transcriptome profiling of whisker follicles in MA self-administered rats. We also investigated alteration in the gene networks (rather than individual gene expression), signalling networks, and the signalling pathways, and the alteration may be responsible for the rewarding effect of MA. The results may serve to reveal the complicated gene-gene interaction mechanisms of the rewarding effect caused by MA.

Results

A MA self-administration rat model. Saline or MA self-administration in rats was performed according to the experimental schedule as indicated in Fig. 1A. Figure 1B shows the number of lever responses during self-administration (2 h each day, 16 days) under the fixed-rate 1 (FR1) schedule. The number of drug infusions in MA self-administration rats gradually increased compared with that in saline self-administration rats and showed <10% variation during the last 3 days of the experiment. The mean number (mean ± SEM) of MA infusions on the final day was 40 ± 7.7. Subsequently, stably responding MA self-administration rats (n = 6) underwent a withdrawal session of 30 days. A two-way analysis of variance (ANOVA) showed a significant difference in the number of infusions between saline and MA self-administered rats (p < 0.01).

RNA-seq and gene expression profiling. RNA-seq was performed using the whisker follicle samples collected from 3 representative rats before self-administration (CON), after MA self-administration (MASA), and after withdrawal (WD) sessions, and the three rats were selected based on the statistical procedure and showed less variation per day during the self-administration session. A total of 8,680 differentially expressed transcripts were analysed using one-way ANOVA in both groups. The quantitative value of each transcript abundance was used to perform a principal component analysis (PCA).

The differentially expressed genes at the three time-points (CON, MASA, and WD) were clustered based on the quantitative results of transcripts following the PCA analysis (Fig. 2A). The differentiation was primarily represented by the first principal component (PC1), accounting for 75.65% of the observed variance. The secondary principal component (PC2) showed 10.79% of the variance. The fundamental difference in RNA abundance between the three time-points accounted for the separation; therefore, principal components could achieve clear separation (Fig. 2A). In the hierarchical clustering analysis (Fig. 2B), it was found that the individual samples were well clustered according to their conditions. Distinct gene expression abundances at each time-point are presented as a heat map in Fig. 2C.

Gene expression pattern analysis. A total of 1,890 genes were identified with same expression patterns after primary data alignment and further filtration (Fig. 3A). In addition, 455 and 205 genes were up- and down-regulated, respectively, at the MASA session compared with the CON session (MASA/CON): 67 and 1,776 genes were significantly up- and down-regulated, respectively, at the WD session compared with the MASA session (WD/MASA) (Fig. 3A). It is noteworthy that more genes (1,776) were down-regulated after MA withdrawal. A time-series correlation of changes in expression identified 666 genes with differential expression patterns in the three rats. According to the mRNA abundance ratios (MASA/CON, WD/MASA), the genes were classified into 6 groups: 3 up-up regulated (Group 1), 422 up-down regulated (Group 2), 2 constant-up regulated (Group 3), 51 constant-down regulated (Group 4), 25 down-up regulated (Group 5), and 163 down-down regulated genes (Group 6) (Fig. 3B,C). Notably, the expression of 422 transcripts in Group 2 gradually increased at the MASA...
session and decreased to the baseline at the WD session (Fig. 3B,C). In contrast, expression of 163 transcripts in Group 6 decreased at the MASA session and then did not decrease further at the WD session (Fig. 3B,C).

Network analysis of differentially expressed genes (DEGs). Physical and functional links of altered genes were investigated based on the assumption that gene pairs that interact or share similar functions tend to interact within the cellular systems. To analyse the changes in mRNA abundances under MA addiction and withdrawal, the functional association network was generated using the 432 seed genes among 666 altered genes based on information about the physical and functional links of these genes. The physical link represents protein-protein interaction, and the functional link represents a relationship between two proteins if they share a substrate in a metabolic pathway and are co-expressed, co-regulated, or involved in the same protein complex. It was found that the network was divided into two prominent subnetworks of up-down regulated (Group 2/red) and down-down regulated genes (Group 6/blue) based on gene expression patterns (Fig. 4A). Interestingly, more links were found within a group (link to the identical group) than between groups (link to the different group). Among 1,466 links in the network, 815 (55.6%) were found to be intra-regulatory links (Fig. 4B). Subsequently, the fraction of links per gene was measured. Links within a group were 2.07-fold more than the expected value (Mann-Whitney U test, \( p \) value = 2.59 e^{-8}; Fig. 4C). The results indicate that the two distinct subnetworks of

Figure 1. Methamphetamine self-administration (MASA). (A) The experimental timeline of saline or methamphetamine self-administration and the sampling time-points of MASA. (B) The number of infusions in saline or methamphetamine self-administration rats. MA self-administered rats (n = 6) had a significantly greater infusion number than did saline self-administration rats (n = 6). Statistical analysis was performed using the two-way analysis of variance (ANOVA) and the student Newman-Keuls test. Error bars represent the mean ± SEM (n = 6). *\( p < 0.05 \), **\( p < 0.01 \), and ***\( p < 0.001 \).

Figure 2. Quality assessment of RNA-sequencing (RNA-seq) data. (A) Grouping of samples via principal component analysis based on gene expression abundance. (B) Hierarchical clustering of genes. (C) A heat-map of the 8,680 expressed genes. Gene expression abundance decreases from red to green.
up-down regulated (Group 2) and down-down regulated (Group 6) genes tend to cluster themselves and might have distinct functional roles in the gene functional network.

**Network mapping of addiction-related genes.** The gene functional network was analysed based on a topological parameter, BC, which reflects the importance of genes in the network. The BC score is the minimum number of links connecting one protein to another in the network; thus, a high BC score implies that genes undergo more number of the shortest paths in the network. Therefore, the protein with the highest BC score derives a lot of importance from its position in the network. To check whether the constructed network contains the gene sets involved in MA addiction, known addiction-related genes from Knowledgebase for Addiction-Related Gene (KARG) database were mapped to the functional network. Subsequently, 1,256 addiction-related genes were collected from 105, 346, 474, and 331 genes mapped from low-throughput (LT) data, mouse, human, and high-throughput (proteomics and microarray) data, respectively, in KARG. As a result, 82 addiction-related genes (6.5%) were mapped in the functional association network that consisted of 432 genes (Fig. 4D). The average BC score of known addiction-related genes was higher than that of other genes (Mann-Whitney U test, \( p = 0.006 \); Fig. 4E). The relative fold-enrichment of known addiction-related genes with the top 10% of BC scores was 2.07-fold higher than that of all genes in the 6 groups (Hypergeometric test, \( p = 5.38 \times 10^{-4} \); Fig. 4F). Finally, top 43 ranked genes, by the BC score, were selected as possible markers of MA addiction (Table 1). Hsp90ab1 (BC score, 0.2127), Akt1 (BC score, 0.1694), and Src (BC score, 0.1171) were identified with the highest BC scores and might act as molecular bridges to connect multiple other proteins in this network. These genes have been previously reported as addiction-related genes\(^{18,29,30}\). Genes with high BC score tend to interact with many different functional groups and are important for controlling information flow in the network. Furthermore, 17 out of 43 genes (39.5%) were mapped as pre-designated addiction-related genes, suggesting that the current approach of network analysis confers a high level of reliability and confidence in the selection of genes that are related to MA addiction. Addiction-related genes with high BC scores could be identified based on their importance in information flow in the altered gene-gene network under addiction-withdrawal conditions.

**Functional analysis using reactome.** To assess the biological relevance of the 43 top ranked genes, functional enrichment analysis was performed using the Reactome database (Reactome, 2004). Biologically related gene sets were identified and labeled with the Reactome terms. The main enriched pathways and genes are
Figure 4. Functional association network analysis of differentially expressed genes. (A) Functional association network in MASA and WD rat whisker follicles. Nodes in the same group are coded with the same colour. Orange and red nodes represent up-regulation in MASA and up-down-regulation in WD, respectively. Yellow and green nodes show constant expression in MASA and up-down-regulation in WD, respectively. Cyan and blue nodes show downregulation in MASA and up-down-regulation in WD, respectively. The links between the same colour nodes are displayed using identical colours. (B) A comparison of link numbers “within” and “between” groups. The number of links according to link type. “Within” is a link to the identical group. “Between” is a link to the different groups. (C) Likelihood scores of “within” groups. Statistical analysis was performed using the Mann-Whitney U test. ***$P < 0.001$. (D) Distribution of addiction-related genes in the network. Links between the same colour nodes are displayed with identical colours. (E) Mean betweenness centrality (BC) scores of the six groups and addiction-related gene group (addiction). Statistical analysis was performed using the Mann-Whitney U test. **$P < 0.01$. (F) Relative fold-enrichment BC scores of the six groups and addiction-related gene group. Statistical analysis was performed using the hypergeometric test. ***$P < 0.001$. 
presented in Table 2. As expected, the pathways included genes that function in the regulation of cell-cell communication, developmental biology, diseases, extracellular matrix organization, hemostasis, the immune system, metabolism, and signal transduction. In addition, several genes associated with the progression of Alzheimer’s disease (AD) were observed (Fig. 5). It is noteworthy that all well-known signalling pathway-related genes, such as transforming growth factor beta 1 (Tgfb1), epidermal growth factor receptor (Egfr), and proto-oncogene tyrosine-protein kinase (Src), activate nuclear factor-kappaB (NF-κB) and thereby regulate the expression of amyloid-β.

| Rank | Genes | Genbank | BC | Addic Re. | Description | Normalised | MASA | WD |
|------|-------|---------|----|----------|-------------|------------|------|----|
| 1    | Hsp90ab1 | NM_001004082 | 0.2127 | O | G4 | Heat shock protein 90 alpha (cytosolic), class B member 1 (Hsp90ab1) | 0.94 | 0.51 |
| 2    | Akt1 | NM_033230 | 0.1694 | O | G2 | V-akt murine thymoma viral oncogene homolog 1 (Akt1) | 1.58 | 0.80 |
| 3    | Src | NM_031977 | 0.1171 | O | G2 | SRC proto-oncogene, non-receptor tyrosine kinase (Src) | 1.31 | 0.66 |
| 4    | Ranbp2 | NM_001191604 | 0.0671 | G5 | | RAN binding protein 2 (Ranbp2) | 0.77 | 1.39 |
| 5    | Mapk14 | NM_031020 | 0.0586 | G4 | Mitogen activated protein kinase 14 (Mapk14) | 0.96 | 0.58 |
| 6    | Egfr | NM_031507 | 0.0568 | G2 | | Epidermal growth factor receptor (Egfr) | 1.30 | 0.67 |
| 7    | Prkacb | XM_00233452 | 0.0447 | O | G2 | Protein kinase, cAMP dependent, catalytic, beta (Prkacb), transcript variant X2 | 1.23 | 0.65 |
| 8    | Gab2l1 | NM_130734 | 0.0436 | G6 | | Guanine nucleotide binding protein (G protein), beta polypeptide 2 like 1 (Gnb2l1) | 0.78 | 0.45 |
| 9    | Cav1 | NM_031556 | 0.0419 | G4 | | Caveolin 1, caveolae protein (Cav1), transcript variant 1 | 3.53 | 0.52 |
| 10   | Cdh1 | NM_031334 | 0.0385 | G4 | | Cadherin 1 (Cdh1) | 1.04 | 0.50 |
| 11   | Cct7 | NM_001191604 | 0.0363 | G6 | | Chaperonin containing Tcp1, subunit 7 (eta) (Cct7) | 0.85 | 0.54 |
| 12   | Jgb1 | NM_017022 | 0.0322 | G2 | | Integrin, beta 1 (Jgb1) | 2.02 | 1.16 |
| 13   | Ppp1ca | NM_031527 | 0.0319 | G6 | | Protein phosphatase 1, catalytic subunit, alpha isoform (Ppp1ca) | 0.81 | 0.50 |
| 14   | Acr1a | NM_001106364 | 0.0313 | G4 | | ARPA1 actin-related protein 1 homolog A, centrinactin alpha (yeast) (Acr1a) | 0.99 | 0.51 |
| 15   | Tgfb1 | NM_021578 | 0.0313 | O | G2 | Transforming growth factor, beta 1 (Tgfb1) | 1.75 | 0.34 |
| 16   | Sec61a1 | NM_199256 | 0.0290 | O | G6 | | Sec61 alpha 1 subunit (S. cerevisiae) (Sec61a1) | 0.73 | 0.31 |
| 17   | Cdk6 | NM_031077 | 0.0274 | G6 | | Cyclin-dependent kinase 6 (Cdk6), transcript variant 2 | 0.85 | 0.51 |
| 18   | Ywhab | NM_019377 | 0.0262 | G6 | | Tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, beta (Ywhab) | 0.76 | 0.52 |
| 19   | Sdc2 | NM_013082 | 0.0250 | G2 | | Syndecan 2 (Sdc2) | 1.77 | 0.67 |
| 20   | Pemd2 | NM_001031639 | 0.0236 | G6 | | Proteasome (prosome, macropain) 26S subunit, non-ATPase 2, (Pemd2) | 0.77 | 0.45 |
| 21   | Jak1 | NM_053466 | 0.0232 | G2 | | Janus kinase 1 (Jak1) | 1.83 | 1.05 |
| 22   | Rela | NM_199267 | 0.0232 | G4 | | V-rel avian reticuloendotheliosis viral oncogene homolog A (Rela) | 1.04 | 0.56 |
| 23   | Vcp | NM_053864 | 0.0218 | G6 | | Valosin-containing protein (Vcp) | 0.78 | 0.51 |
| 24   | Srebf2 | NM_001033694 | 0.0211 | G6 | | Sterol regulatory element binding transcription factor 2 (Srebf2) | 0.94 | 0.55 |
| 25   | Apolipoprotein | NM_014593 | 0.0206 | G6 | | Apolipoprotein (Apolipoprotein) | 0.79 | 0.44 |
| 26   | Npl1 | NM_145098 | 0.0201 | G2 | | Neuritin 1 (Npl1) | 1.87 | 0.63 |
| 27   | App | NM_019288 | 0.0197 | G2 | | Amyloid beta (Aβ) precursor protein (App) | 2.71 | 0.83 |
| 28   | Eef1g | NM_001004223 | 0.0197 | G6 | | Eukaryotic translation elongation factor 1 gamma (Eef1g) | 0.77 | 0.33 |
| 29   | Stat5a | NM_017064 | 0.0191 | G2 | | Signal transducer and activator of Transcription 5A (Stat5a) | 2.18 | 0.64 |
| 30   | Itch | NM_001005887 | 0.0187 | G2 | | Ichi E3 ubiquitin protein ligase (Itch) | 1.22 | 0.87 |
| 31   | Atg10 | NM_012504 | 0.0183 | G2 | | ATGase, Na+/K+ transporting, alpha polypeptide (Atg10) | 1.83 | 0.63 |
| 32   | Zmir1 | NM_001083933 | 0.0176 | G2 | | Zinc finger, MIZ-type containing 1 (Zmir1) | 1.72 | 0.50 |
| 33   | Glil | NM_00191910 | 0.0161 | G2 | | GLI family zinc finger 1 (Glil) | 3.78 | 1.42 |
| 34   | Prkcg | NM_013012 | 0.0158 | G6 | | Protein kinase, cGMP-dependent, type II (Prkcg) | 0.51 | 0.09 |
| 35   | Cbp2 | NM_053335 | 0.0153 | G2 | | C-terminal binding protein 2 (Cbp2) | 1.24 | 0.92 |
| 36   | Hdh | NM_133618 | 0.0147 | G6 | | Hydroxyacyl-CoA dehydrogenase/3-ketoacyl-CoA thiolase/enoyl-CoA hydratase (trifunctional protein), beta subunit (Hdh) | 0.69 | 0.48 |
| 37   | Nflk2 | NM_001008349 | 0.0144 | G2 | | Nuclear factor of kappa light polypeptide gene enhancer in B-cells 2, p49/p100 (Nflk2) | 1.57 | 0.85 |
| 38   | Stk11 | NM_00108069 | 0.0143 | G6 | | Serine/threonine kinase 11 (Stk11) | 0.82 | 0.43 |
| 39   | Vcl | NM_00107248 | 0.0141 | G2 | | Vinculin (Vcl) | 1.73 | 0.73 |
| 40   | Pip4k2b | NM_053550 | 0.0136 | G2 | | Phosphatidylinositol 3-phosphate 4-kinase, type II, beta (Pip4k2b) | 1.46 | 0.74 |
| 41   | Tie3 | NM_053400 | 0.0134 | G2 | | Transducin-like enhancer of split 3 (Tie3) | 0.70 | 0.35 |
| 42   | Map3k3 | NM_01107058 | 0.0133 | G2 | | Mitogen activated protein kinase (Map3k3) | 1.41 | 0.72 |
| 43   | Fbxw11 | NM_00106993 | 0.0129 | G2 | | F-box and WD repeat domain containing 11 (Fbxw11) | 1.16 | 0.68 |

Table 1. List of genes with top 10% betweenness centrality (BC) scores in the network.
precursor protein (APP), which is associated with AD. Moreover, most genes associated with biosynthesis of heparan sulfate, a member of the glycosaminoglycan family, were upregulated at the MASA session (Fig. 5).

**Discussion**

MA abuse is a major health concern and a social problem worldwide, and the mechanism of addiction still remains unclear. Clinically, it is crucial to understand MA addiction because of a current lack of effective pharmacotherapy and a high recurrence rate. Therefore, it is important to make an accurate diagnosis of MA abuse and addiction in clinical medicine. In fact, non-invasive diagnostic tools or reliable biomarkers used to determine MA abuse and addiction are being sought.

The purpose of the present study was to investigate the transcriptional changes caused by rewarding effects in MA self-administration rats. Drug addiction is not a static condition but a chronic relapsing disorder, which develops with the repetition of rewarding and withdrawal effects caused by temporary drug use and abstinence, respectively. Therefore, it is not enough to only study static changes of genes for MA addiction. The transcriptional gene changes caused by initial reward and withdrawal could affect the following stages and play a key role in MA addiction. To find diagnostic genes in this process, we examined dynamic changes in mRNA expression in the rat whisker follicle during MA self-administration. We used the short-access rat model of MA self-administration and measured the number of drug infusions that each rat chose (reward rate), and it was found that the rats showed an increased rewarding effect upon repeated MA administration. Analysis of RNA-seq data revealed that time-dependent changes of gene expression patterns at the three time-points (CON, MASA, and WD) were different. Notably, the majority of genes showed a similar pattern of changes in expression: an initial increase followed by a decrease during MASA and WD, suggesting that these genes may play a similar role in maintaining the reward state caused by MA.

Network analysis is a tool for gene prioritization and can facilitate a better understanding of network interconnections among interactive genes. Functional gene-gene association network analysis is an effective way to understand the interaction of a large number of genes. The importance of genes can be determined by measuring BC scores in the network. In this study, a topological centrality analysis was performed using a functional association network to narrow down the candidate genes related to MA addiction, and 43 genes were selected as core genes in the network.

The top three core genes were heat shock protein (HSP) 90-beta 1 (Hsp90ab1), RAC-alpha serine/threonine-protein kinase (Akt1), and Src. They have been previously reported as drug addiction-related genes; therefore, it is possible that they play an important role in MA addiction. The gene with the highest BC score was Hsp90ab1 (BC score, 0.2127), and Hsp90-beta protein, encoded by Hsp90ab1 gene, is a molecular chaperone maintaining the functional stability and viability of cells. The expression of Hsp90ab1 in the rat frontal cortex increases after morphine self-administration. Furthermore, Akt1, which had the second highest BC score (0.1694) in this study, is implicated in excessive alcohol drinking behaviours. Src-family kinase activation is necessary for incentive, motivational, and/or memory processes that promote contextual cocaine-seeking behaviours. To determine the biological function of 43 genes that could act as a functional core in the constructed gene network, we performed functional enrichment analysis using the Reactome database. Reactome categorises large quantities of genes and proteins into functions and pathways based on a massive database.

Using the Reactome database, we identified the main enriched pathways of the selected 43 genes and discovered eight functional groups: cell-cell communication, developmental biology, diseases, extracellular matrix organisation, hemostasis, the immune system, metabolism, and signal transduction. Notably, a prominent increase in the expression of genes related with AD and heparan sulfate biosynthesis was found. AD is a chronic neurodegenerative disease and the most common age-related dementia. Previous studies suggest that MA abuse is related to long-term damage in the human brain. In fact, clinical studies using proton magnetic resonance spectroscopy showed that the level of N-acetyl-aspartate, a neuronal marker, reduced by 5% in MA abusers, and the reduction was also observed in several brain diseases, such as AD, stroke, and epilepsy. The representative characteristics of AD are progressive loss of cognitive abilities, severe neurodegeneration, and neuroinflammation. One of the hallmarks of AD is the formation of neuritic plaques, which contain filamentous extracellular β-amyloid (Aβ) aggregates. Aβ is generated by sequential cleavage of APP by NF-κB-dependent activation of β- and γ-secretase. A previous study revealed that MA enhanced Aβ accumulation through alpha7 nicotinic acetylcholine receptor in brain endothelial cells. In agreement with the aforementioned results, an increased expression of NF-κB, APP, and Aβ was found in the whisker follicles in MA self-administered rats in this study.

Heparan sulfate biosynthesis is another interesting pathway in this study. Heparan sulfate proteoglycans (HSPGs) are macromolecules located in the cell surface and extracellular matrix. HSPGs consist of core protein-linked glycosaminoglycan chains. The core proteins of HSPGs are agrin, perlecans, glypican, and syndecans (Sdc). In this study, a significant increase in the expression of Sdc2 was found at the MASA session. In addition, most of genes included in the heparan sulfate pathway showed the same expression pattern as did Sdc2. According to previous studies, Sdc has four categories of functions: (1) growth factor receptor activation; (2) matrix adhesion; (3) cell to cell adhesion; and (4) tumour suppression and progression. Sdc2 is involved in synaptic formation and Sdc3 has been identified as a resilience factor of cocaine addiction. Consequently, transcriptional alteration of Sdc2 at the MASA session may be an important marker of MA addiction.

Drug addiction is a complicated, chronic disorder and is conceptualised as a cycle comprising rewarding effect, withdrawal effect, craving, and relapse to drug seeking behaviour. Briefly, intake of drugs initially causes rewarding effect, and tolerance and withdrawal effects gradually occur after repeated use of the drugs. To escape the effects, craving to the drug intake arises, leading to relapse of drug-seeking behaviour. Therefore, a validated model of intravenous drug self-administration relevant to the transition to addiction is needed to uncover the
| Pathways                          | Genes               | p-Value |
|----------------------------------|---------------------|---------|
| **Cell-cell communication**      |                     |         |
| Cell junction organization       | Vcl                 | 0.02    |
| Cell-cell junction organization  |                     |         |
| Adherens junctions interactions  | Vcl                 | 0.01    |
| **Development biology**          |                     |         |
| Axon guidance                    |                     |         |
| L1CAM interactions               |                     |         |
| CHL1 interactions                | Nrp1, Itgb1         | 0.01    |
| **Disease**                      |                     |         |
| Diseases of signalling transduction |                 |         |
| Signalling by TGF-beta Receptor Complex in Cancer | | |
| Loss of function of TGRBR1 in Cancer | | |
| TGFBR1 LBD Mutants in Cancer     | Tgfb1               | 0.05    |
| Loss of function of TGRBR2 in Cancer |               | 0.03    |
| TGFBR2 MSI Frameshift Mutants in Cancer | | |
| Signalling by WNT in cancer      |                     |         |
| TCF7L2 mutants-unbound CTBP       | Ctbp2               | 0.04    |
| Oncogenic MAPK signalling        |                     |         |
| Signalling by high-kinase activity BRAF mutants | Src, Ywhab, and Vcl | 0.03    |
| Signalling by moderate kinase activity BRAF mutants | Src, Ywhab, and Vcl | 0.05    |
| Paradoxical activation of RAF signalling by kinase inactive BRAF | Src, Ywhab, and Vcl | 0.05    |
| **Diseases of Immune System**    |                     |         |
| Diseases associated with the TLR signalling cascade | | |
| IKBKG deficiency-caused anhidrotic ectodermal dysplasia with immunodeficiency (EDA-ID) (via TLR) | Nfkb2, Rela | 0.01 |
| IkB variant leads to EDA-ID     | Nfkb2, Rela         | 0.01    |
| **Extracellular matrix organization** |                 |         |
| Non-integrin membrane-ECM interactions | | |
| Syndecan interactions            | Itgb1, Sdc2         | 0.04    |
| **Hemostasis**                   |                     |         |
| Cell surface interactions at the vasculat wall | | |
| Basigin interactions             | Itgb1, cav1         | 0.01    |
| **Immune System**                |                     |         |
| Adaptive immune system           |                     |         |
| Costimulation by the CD28 family |                     |         |
| CTLA4 inhibitory signalling      | Src and Akt1        | 0.05    |
| Innate Immune System             |                     |         |
| DDX58/IFIH1-mediated induction of interferon-alpha/beta | | |
| TRAF6 mediated NF-kB activation  | App, Nfkb2, and Rela | 0.02 |
| Cytosolic sensors of pathogen-associated DNA | | |
| ZBP1 (DAI) mediated induction of type 1 IFNs | | |
| RIP-mediated NFkB activation via ZBP1 | App, Nfkb2, and Rela | 0.03 |
| DEx/H-box helicases-activated type I IFN and inflammatory cytokines production | App, Nfkb2, and Rela | 0.03 |
| C-type lectin receptors (CLRs)   |                     |         |
| CLEC7A (Dectin-1) signalling     |                     |         |
| Dectin-1 mediated noncanonical NF-kB signalling | Psmd2, Nfkb2, Fbxw11, and Rela | 0.03 |
| Cytokine Signalling in Immune system | | |
| Signalling by Interleukins        |                     |         |
| Interleukin-1 family signalling  |                     |         |
| Interleukin-1 processing         | Nfkb2 and Rela      | 0.02    |
| **Metabolism**                   |                     |         |
| Metabolism of lipids             |                     |         |
| Triglyceride metabolism          |                     |         |
| Triglyceride catabolism          |                     |         |
| Phospholipid metabolism          |                     |         |
| PI Metabolism                    |                     |         |
| Continued                        |                     |         |
complicated clinical situation in human addicts. In the present study, rats were allowed to self-administer MA 2 h daily for 16 days to determine the mechanism underlying the transition from initial drug use to drug addiction. However, the present study has some limitations. For example, a short access rat model of MA self-administration was used in this study. Recent studies have shown that rats with extended access to MA, compared with rats with short access to MA, showed enhanced MA-primed reinstatement of drug-seeking and cognitive deficits52–55. Moreover, extended limited daily access to MA self-administration in animal models mimics the course of MA administration in human55. Therefore, the model in the present study can be used to study the reward process, which is an initial step of the MA addiction. However, a follow-up study with long access self-administration would be needed to understand MA addiction in human.

In conclusion, the present findings suggest that the alteration of the gene expression and signalling networks in the whisker follicles in MA self-administered rats can be used to understand the biological changes and MA rewarding effects induced by MA self-administration. Further studies should be needed to determine the differential changes in gene expression between the whisker follicle and the brain to strengthen our conclusion.

### Methods

#### Animals.
Adult male Sprague-Dawley rats (Daehan Animal, Seoul, Republic of Korea), 310–350 g in weight, were housed individually in the laboratory animal facility in room temperature (22 ± 2 °C) and humidity (60 ± 2%) controlled cages on a 12 h light/dark cycle (lights on at 7:00 AM). Animals had access to food and water ad libitum. All experimental procedures were approved by the institutional animal care and use committee at Daegu Hanny University, Daegu, Republic of Korea (DHUMC-D-13002-PRO-02, 2013). The experiments were carried out in accordance with the university's scientific research guidelines and regulations.

#### MA self-administration and sample collection.
MA self-administration was conducted as previously described56. Briefly, rats were habituated to their environment for 1 week, and they were then implanted with chronic indwelling jugular catheters and assigned to the control (n = 6) or MA self-administration (n = 6) group. One week after surgery, rats were given food training. Subsequently, they were treated with a FR1 schedule of MA self-administration with 2 h access per day to MA hydrochloride (0.05 mg·kg·infusion⁻²) dissolved in sterile heparinised saline (3 U·mL⁻¹) for 2 weeks. If animals had shown the stable infusion numbers for 3 consecutive days (variation less than 10% of the mean infusion number of the 3 days), the mean infusion number was regarded as the baseline. Saline self-administered rats received heparinised saline infusions during the self-administration period. All responses were recorded using MED associated software. Samples were collected 3 times during the experiment: Control (CON) samples were collected before MASA; MASA samples were collected within 2 hours after the last MA self-administration; and WD samples were collected 1 month after MASA.

#### RNA isolation.
Whisker samples were collected and stored at −80 °C before RNA extraction after rats were sacrificed. Total RNA was isolated from the whisker roots (hair follicles) using the spin column type RNA extraction kit (Bioneer, Republic of Korea). The quality and concentration of RNA were measured using a filter-based multi-mode microplate reader (FLUOstar Omega, BMG LABTECH, Germany). Residual genomic DNA was removed by incubation with RNase-free DNase I for 30 min at 37°C.

#### Construction of transcriptome libraries.
The mRNA sequencing libraries were prepared according to the Illumina TruSeq RNA sample preparation kit v2 (Illumina, Inc., San Diego, CA, USA). Briefly, 1 μg total RNA was used for polyA mRNA selection using streptavidin-coated magnetic beads, followed by thermal fragmentation of selected mRNA. The 200–500 bp fragmented mRNA was used as a template for cDNA synthesis by reverse transcription.
transcriptase with random primers. The cDNA was converted into double-stranded DNA that was end-repaired (to incorporate the specific index adaptors for multiplexing), purified, and amplified for 15 cycles. The quality and functionality of the final amplified libraries were examined with a DNA high sensitivity chip on an Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, CA, USA). Libraries were prepared according to the Illumina's protocol. Briefly, random fragmentation of the DNA or cDNA, followed by 5′ and 3′ adapter ligation, was conducted. Alternatively, “tagmentation” combined the fragmentation and ligation reactions into a single step, which greatly increased the efficiency of the library preparation process. Subsequently, adapter-ligated fragments were PCR amplified and gel-purified. The concentration of each library was measured by real-time PCR. Agilent 2100 Bioanalyzer was used for profiling the distribution of insert size. For cluster generation, the library was loaded into a flow cell where fragments were captured on a lawn of surface-bound oligos complementary to the library adapters. Each fragment was then amplified into distinct clonal clusters through bridge amplification. After cluster generation was completed, the clustered libraries were sequenced by Illumina HiSeqTM 2500 according to the manufacturer's instructions (Illumina Inc., USA), and paired-end sequencing was performed for 100 cycles using TruSeq rapid SBS kit or TruSeq SBS kit v4. The sequencing protocol, the HiSeqTM 2500 system user guide part #15011190 Rev. V HCS 2.2.58, was used in this study.

**Sequencing data analysis.** Sequencing raw images were generated using HiSeqTM Control Software (HCS v2.2), and base calling was performed through an integrated primary analysis software called RTA (v1.18). Images generated by HiSeqTM2500 were converted into nucleotide sequences by base calling and were stored in the FASTQ format using the Illumina package bcl2fastq (v1.8.4). Clean reads were generated by filtering the dirty reads, which contained adaptors and unknown or low phred quality scored-bases, from raw reads. Clean reads were mapped to reference Rattus norvegicus genome (rn6) and gene sequences using a conventional Tuxedo protocol. Reading mismatches of ≤5 bases were allowed in the alignment. Reads that matched with reference rRNA sequences were also mapped and removed. TopHat2 software and the Cufflinks package (v2.2.1) were used to calculate FPKM (Fragments per kb per million reads) values for each gene used for subsequent data analysis. DEGs were analysed using one-way ANOVA, Student’s t-test (p < 0.05), and fold changes. Hierarchical clustering and PCA were also performed with the GeneSpring 7.3 software (Agilent Technologies).

**Gene expression pattern analysis.** Transcripts that were differentially expressed between the groups were analysed using one-way ANOVA. After primary data alignment and further filtration to find transcripts with >1.1 or <0.9 fold-change between groups, 1,890 transcripts were quantified, and 666 genes were next identified with same patterns in both MASA/CON and WD/MASA.

**Functional association network analysis.** The STRING-database (ver. 10.0) was used to conduct a rat gene functional association network using physical and functional link information extracted via the retrieval of interacting with a confidence score over 400. Network visualization and calculation of BC were performed using Cytoscape 2.8.
Statistical analysis. Data were analysed using a two-way ANOVA followed by a Student Newman-Keuls test in the self-administration experiment. Data from the self-administration group were compared with those from the saline self-administration group, and *p < 0.05, **p < 0.01, or ***p < 0.001 was considered statistical significance. The differences in the RNA-seq results within groups were evaluated by PCA. Differentially expressed transcripts between the groups were calculated using a one-way ANOVA. The Mann-Whitney U test was performed to compare links in the network, and the hypergeometric test was used for the comparison of relative fold-enrichment in the network.

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**Acknowledgements**

This study was supported and funded by the Basic Science Research Program of the National Research Foundation of Korea (NRF), the Ministry of Education (NRF-2016R1A6A1A03011325) and the Bio & Medical Technology Development Program of the NRF, and the Ministry of Science, and ICT & Future Planning (NRF-2015M3A9E1028327).

**Author Contributions**

S.H.S. and W.J.J. performed the experiments and participated in writing the manuscript. J.H. performed the network analysis and wrote the manuscript. B.P., J.H.J., Y.H.S., and C.H.Y. participated in the experimental design and analysed the results. S.L. and C.H.J. were responsible for the study concept and design and supervised the manuscript editing.

**Additional Information**

**Competing Interests:** The authors declare no competing interests.

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