Placebo effects and the molecular biological components involved

Lei Cai, Lin He

ABSTRACT
Pharmacologically inactive substances have been used in medicine for more than 700 years and can trigger beneficial responses in the human body, which is referred to as the placebo effects or placebo responses. This effect is robust enough to influence psychosocial and physiological responses to the placebo and to active treatments in many settings, which has led to increased interest from researchers. In this article, we summarise the history of placebo, the characteristics of placebo effects and recent advancements reported from the studies on placebo effects and highlight placebo studies to identify various molecular biological components associated with placebo effects. Although placebos have a long history, the placebo concept is still in its infancy. Although behavioural, neurobiological and genetic studies have identified that molecules in the dopamine, opioid, serotonin and endocannabinoid systems might be targets of the placebo effect, placebo studies with a no-treatment control (NTC) are necessary to identify whole-genome genetic targets. Although bioinformatics analysis has identified the molecular placebome module, placebo studies with NTCs are also required to validate the related findings.

INTRODUCTION
In the 1300s, the word ‘placebo’ first appeared in a Latin translation of the Hebrew Bible, with the original meaning of ‘to walk’. Later, because of a mistranslation, placebo was defined as ‘to please’. With this positive meaning, the word placebo was recorded as meaning ‘to please’ in early usage. In the mid-to-late 1700s, the term placebo began to be used as medical jargon. The prevailing opinion is that an English physician and pharmacologist, William Cullen, introduced the concept of placebo in his clinical lectures given in 1772, but another opinion is that before him, another British physician, Alexander Sutherland, who was familiar with the water cure, first used the word placebo in his book and tried to revive some ancient medical doctrines. At that time, according to William Cullen’s introduction, the word placebo referred to drugs that were administered to satisfy a patient’s desire for a remedy, however the physician thought that the placebo was ineffective for treatment of severe diseases.

In this article, we operationally define placebo as an inert treatment, such as: drugs or surgery, which can be used to simulate administration of a real medical intervention and has been used as an indispensable control in randomised clinical trials (RCT). In 1784, Benjamin Franklin and Antoine Lavoisier used placebos as controls in a trial in which they exposed patients to so-called ‘animal magnetism’ objects or normal objects without identifying them and found that the patients’ responses were similar. In RCTs, placebos are used as a methodological tool to challenge and debunk ineffective treatments and have become a mainstream of modern medicine. RCTs are the gold standard for tests of safety and efficacy of novel medical treatments and include definite steps, such as double-blind randomisation, besides the use of placebo. The ultimate goal of medical treatment is to heal (ie, to control or cure an illness) and provide symptom relief or comfort. However, at some point in the progression of diseases, there are no cures available or methods to ease the suffering. In such situations, empathic healthcare can predispose patients to shifts in the perceptions of their body status, cause decreased reactivity to the underlying pathophysiology and relieve unnecessary suffering. In other words, patients who receive more empathic healthcare may have more hope and, consequently, relief. Interestingly, expectations or hopes can trigger bodily responses, particularly during childhood. Thus, placebos are necessary in RCTs and have some psychotherapeutic value.

Compared with well-targeted very efficacious medications, placebos should give only modest results in RCTs. An issue in RCTs is that controlled patients are not fully informed of whether they are specifically receiving a placebo treatment, so trials with placebo as the control sometimes pose an ethical dilemma if the test drug is effective. However, a placebo itself can indicate whether the...
efficacy of a new medical intervention or treatment is sufficient to justify its use. Notably, in the 1950s, Beecher performed a meta-analysis by combining the placebo treatment data collected from 15 studies on different diseases and found that placebos led to an approximately 35% improvement in symptoms. In some RCTs, placebos without any known active principles are of more help in relieving some patients’ suffering than are tested medical treatments. The phenomenon of improvement of the symptoms of a patient who has taken a placebo in a RCT is called placebo effect or placebo response. The placebo effect is so robust that it can influence psychological and physiological responses to the placebo and to subsequent active treatments in many settings, so it has attracted increasing attention of researchers.

There are several theories that attempt to explain the mechanism of the placebo effect: the expectancy theory, classical conditioning accounts, context effects and the meaning response. Recently, a proposed framework based on integrative framework theory by Colloca and Miller has been widely accepted. The integrative framework theory emphasizes that cues of a different nature (ie, verbal, contextual, social) may be integrated to generate key treatment expectancies, which can influence the effects of active or placebo treatment. Thus, empirical findings for placebo can be integrated into a single conceptual model rather than other complex dual mechanisms.

In this article, on the basis of the history of placebo, we summarise the characteristics of placebo effects and the recent advancements reported from studies on placebo effects, as well as highlight placebome studies to identify various molecular biological components of placebo effects.

CHARACTERISTICS OF PLACEBO EFFECTS
Placebo effects are actually the body’s responses to a general expectancy through absorbing some cues, including physical and psychological ones. However, some major factors identified to affect placebo effects are patients’ reporting bias, regression to the mean and the physiological variation of illnesses in RCTs. Since a patient may tend to report improvement that has not actually occurred under some circumstances, it is easy to treat reporting bias as a true effect of placebo based on subjective outcomes. The regression to the mean is a statistical phenomenon in which a variable is extreme on its first measurement, but with increases in the number of measurements, the variable’s value approaches the mean or average, that is, the placebo effect is high on the first measurement but may be lower on the second and subsequent measurements. Since the pathological conditions of many diseases undergo natural changes, some patients may have spontaneous remission along with the natural waxing and waning of an illness without any treatment or intervention. Thus, familiarity with the characteristics of placebo effects will provide a better understanding of how they work.

Placebo effects usually have the following characteristics: (1) since placebos have no inherent therapeutic power, they rarely cure the illness but may provide relief of some patients’ subjective symptoms, such as pain; (2) placebo effects widely vary in patients with different diseases and in patients with the same disease treated with different medicines; (3) there are also adverse consequences of placebo effects, that is, the so-called ‘nocebo effect’. Up to 26% of patients randomly assigned to placebos in RCTs are estimated to discontinue the use of placebos because these patients have perceived adverse effects. Actually, the psychosocial factors that induce nocebo effects can also cause adverse medication effects. Placebo effects are beyond the reach of medical intervention or treatment, the patients’ cognitive level on the treatment and/or the physician–patient relationship can enhance the effectiveness of medical treatment. An interesting study demonstrated that the patients who took the real drug rizatriptan but labelled as ‘placebo’ had no different outcomes from those taking placebos labelled with ‘rizatriptan’. But when patients took the real drug rizatriptan correctly labelled as ‘rizatriptan’, the effect of this drug increased by about 50%. Another study also obtained similar results in which the effects of open versus hidden administration of morphine for postoperative pain, beta-blockers for cardiovascular function, subthalamic stimulation for Parkinson’s disease and diazepam for anxiety were compared and open treatment was found to induce significantly greater improvement than that of the hidden one.

In the view of Miller and Colloca, the placebo effect is a learnt response generated by expectancies via the central nervous system. When a patient has an active or placebo treatment, the different cues may cause the patient to remember the previously experienced sensations and thereby develop an expectancy. Different cues can converge into a single conceptual model to generate key expectancies, which is a more general state that relates to consciousness or subconsciousness according to the specific process involved. To understand both placebo and nocebo effects, although they have different psychological mechanisms, the general conceptual framework is considered to be the same as the expectancies determined by prior experience. Increasing evidence supports the ideas that the placebo is not limited to inert agents but many active treatments may also have the similar effects of placebo. It is necessary to consider the placebo effect when carrying out any medical treatments.

PLACEBO EFFECT STUDIES
In the past, behavioural instruments were used to study the mechanism of placebo effects. Overall, behavioural studies suggest an important role of learning in the placebo effect, including individual training and social learning. It has been found that the individual training would ...

duration and method (continuous or partial reinforce-
ment) may influence the results of various placebo effects, the verbal and social cues may influence the results of training and all combined available factors present during the clinical treatment may determine the overall results of placebo effects. Thus, there has been limited success for this approach because these instruments cannot explain the complex placebo response states that shift based on a patient’s beliefs, expectations and previous experiences.

With advances in neuroimaging, we have explored a number of neurobiological mechanisms of the placebo effect. Through this technique, placebo effects have been shown to be biological responses to psychosocial cues associated with the medical treatments that rely on complex neurotransmitters involved in neurobiological mechanisms, such as cannabinoids and dopamine, and on some brain regions, such as amygdala, anterior insula and prefrontal cortex in placebo analgesia. Although objective neurobiological pathways have been revealed to correlate with placebo effects, no evidence supports that placebo effects can alter the pathophysiology of disease. These substantial advancements in the placebo effect are essential for evaluating drug effects.

To facilitate pharmaceutical development, rigorously characterised placebo effects based on each patient may be of great value to patients and researchers. For drug development, an underlying goal of RCTs is to find a difference between active treatment and the placebo control. Knowing likely placebo responders could improve trial designs to detect such a difference, and modify treatment approaches by allowing for more efficient medication dosages. A pressing issue in treatment is to solve the conflict of the disclosure of drugs’ adverse effects with the avoidance inducing nocebo effects. Addressing this issue depends on characterised placebo effects based on each participant without deception. A striking finding in RCTs is the effect of possible placebo pathway genes on the treatment with both the placebo and drug, which has demonstrated that some drugs have placebo–drug interactions as a result of shared molecular targets. Thus, precise knowledge of molecular biological components of placebo effects promises to lead to greater understanding of the underlying mechanisms, although the environmental factors that surround a patient make it an ongoing challenge.

It has been widely accepted that clinical outcomes are affected by the interplay of genetic factors and environmental factors. As is known, the placebo effect is a complex phenotype affected by a subject’s beliefs, expectations and previous experiences. Additionally, placebo effects were reported to have been observed in 77.5% of subjects in naltrexone trials for the treatment of alcoholism and there are other types of placebo effects that have been observed in patients with certain symptoms of pain, headache and nausea. Since an individual’s genetic make-up is considered to be a stable inner trait, genetic variation is an important factor that influences placebo effects. Genetic variations can lead to function abnormality of genes, RNA and protein networks and form an individual’s genetic response characteristics. Thus, greater understanding of genetic impacts on the placebo effect may help distinguish active treatment effects with placebo effects in certain research designs and help obtain the precise knowledge of the molecular biological components of specific placebo effects.

The recent availability of large-scale -omics data of genes, RNA and proteins (ie, genomics, epigenomics, transcriptomics, proteomics, and so on) offers a potential new approach to identifying molecular targets for placebo effects. The recently emerging concept of a placebome was proposed through collecting -omics data, such as genomic data, to unpack influences of genome-derived molecules on placebo effects. Since a better understanding of inner genetic influences of placebo effect is critical for evaluating and maximising the efficacy of medical treatment, knowledge of the placebome is of potential benefit to develop novel strategies for clinical trial designs, reduce trial cost and improve the understanding of the mechanisms underlying the placebo effect. Thus, placebome studies are justified for these reasons.

Despite the promise of placebome to discover and develop more effective personalised medicine, it is probably worth noting that, while still in a relatively nascent phase, -omics studies in psychiatry have yielded little tangible benefit to date regarding predictive therapeutic benefits (other than identifying individuals at risk for side effects due to variations in drug metabolism). It is also true that no genome-wide association studies (GWAS) related to placebo effects have yet been conducted. Thus, the search for genomic targets associated with placebo effects is in its infancy. In fact, many placebo-controlled RCTs have used genomics data, but all lacked a no treatment control (NTC), which is usually used to distinguish genuine placebo effects from regression to the mean and natural changes in an illness. Furthermore, in RCTs, addressing more ethical issues often takes precedence over use of an NTC since use of a placebo is thought to be treatment, but placebo employment without the patients’ knowledge may violate their rights of informed consent and cause patients to distrust their doctors. In a recent placebome study without NTC, placebo effects varied between 25% and 75% in psychopharmacology, and because of the variable placebo effects, about 50% of antipsychotic clinical trials are not found to support the superiority of tested drugs over placebo, although the identified placebome module may be significantly similar to the depression and anxiety modules in the human interactome. Thus, inclusion of an NTC in studies to investigate the placebome is required in future studies.
known cues that trigger placebo effects may be similar but not identical to previously experienced cues. Thus, there may be multiple mechanisms for placebo effects and there exists a key mechanism that has not been identified yet. In placebo analgesic studies, the learning mechanisms have been identified to affect neural and cognitive aspects of the placebo effects. In nocebo hyperalgesic studies, two regions, the rostral anterior cingulate cortex (rACC) and the periaqueductal grey (PAG), have been found to facilitate expectation-induced pain. Neuroimaging has shown that the neural interactions between the prefrontal areas, brainstem and spinal cord can mediate the nocebo effect. Through modulation of connecting within the rACC-PAG spinal axis, cognition interacts with the pain pathway to modulate pain and nociceptive processing at the spinal level. Further investigations of the molecular biological components that have yielded the behavioural and neuroimaging data are needed to elucidate the mechanisms of placebo or nocebo effects in greater depth.

The nocebo may consist of multiple intersecting pathways, and there may be genetic overlap between placebo, disease, and treatment; specifically, the genes possibly involved in the placebo pathway exert effects in the drug pathway or disease pathogenic pathway. Based on previous studies of disorder treatments, the genes of placebo effects may have wide effects on dopamine, opioid, endocannabinoid and serotonin signalling pathways. These four signalling pathways have been identified to affect neural and cognitive aspects of the placebo effect and are viewed as important processes in the subjective experience of symptom relief related to the placebo effect. In the next section, we summarise the molecular biological components involved in these signalling pathways.

Molecular Biological Components of Placebo Effects

The importance of identifying molecular biological components of the placebo effects is not limited to excluding the most likely placebo responders in RCTs to maximise the efficacy of medical treatments. Purposefully inhibiting the placebo effects by a drug in advance could minimise the placebo effect and interfere with evaluation of the effect of medical treatments in an objective manner. Through integrating the behavioural, neurobiological and genetic findings on the placebo effect, the dopamine, opioid, endocannabinoid and serotonin signalling pathways are used as the primary means for identifying molecular biological components through analysis of the genetic variants. However, it is worth noting that besides these four systems, various psychological and biological factors across different psychiatric diseases might also mediate placebo effects, but these have not been studied yet. Along with advances in knowledge about the neurotransmitters and neural pathways, increasingly specific candidate genes influencing the placebo effect have attracted greater attention. In particular, in the past years, high-throughput analysis technologies have produced a large number of gene and protein–protein interaction data that have stimulated studies of systems biology. These -omics data provide unprecedented opportunities to investigate the molecular targets of placebo effects at the systems level by conducting placeboome studies. As mentioned earlier, certain genes or gene products may mediate placebo effects in individual patients together, and there is potential molecular overlap among placebo and medical treatment effects and the disease, which highlights the complexity of placebo studies and the importance of identifying the molecular biological components of placebo effects.

Table 1 summarises the genes with possible involvement in placebo effects, although multifaceted and complicated nature of placebo effects should be considered when reviewing the table. It is important to be aware that it would be unrealistic to expect several genetic variants alone to influence a majority of placebo effects because there is a lot of evidence supporting multiple mechanisms. This idea is also supported by a number of GWAS, which have demonstrated that almost all common variants affect complex traits with very small effect sizes. The genetic association studies require a certain number of samples to balance type I errors and power. Assessing multiple variants in one experiment may increase type I errors without any controls for multiple comparisons. Additionally, a significantly larger sample size is required to increase the power that may be reduced by controlling for multiple comparisons. Thus, the previous genetic studies of placebo effects with a relatively small number of participants may have underestimated and overestimated the role of some variants. To balance power and type I errors, a study design using small twin or sibling samples to investigate the genetic contribution to placebo effects is plausible to increase power and reduce noise. This approach could obtain the highly similar genetic background, and therefore clearly identify the possible genes involved in placebo effects. An increasing number of studies have demonstrated that the neurotransmitter and neurological pathways can mediate placebo effects, and have provided candidate genes for further studies. First, a placebo was found to induce the pain suppression system of the body, which can be blocked by an opioid receptor antagonist. This finding demonstrated that the opioid signalling pathways may be involved in a placebo analgesic effect. Furthermore, activation of some brain regions induced by expectation of analgesia is related to endogenous opioid transmission and analgesia. Further physiological experiments have demonstrated that the endocannabinoid signalling pathway is also implicated in the placebo analgesia. Based on the finding of analgesic effects of opioid receptor signalling, expectancy of reward is postulated as a key general contributor to the placebo effect. In a pain model, anticipation of the placebo effect stimulated the activation of opioid and dopamine receptor in brain, and higher placebo effects have been found to correspond to higher levels of dopamine receptor activation. Additionally, both
Table 1  Summary of genetic targets of placebo effects

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Gene symbol</th>
<th>Location</th>
<th>SNP</th>
<th>Placebo pathway</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Catechol-O-methyltransferase</td>
<td>COMT</td>
<td>22q11.2</td>
<td>rs4680, rs4633</td>
<td>Dopaminergic, serotoninergic</td>
<td>51</td>
</tr>
<tr>
<td>Monoamine oxidase</td>
<td>MAOA</td>
<td>Xp11.3</td>
<td>rs6323, rs6609257</td>
<td>Dopaminergic, serotoninergic</td>
<td>53, 58</td>
</tr>
<tr>
<td>Nuclear receptor subfamily 3 group C member 1</td>
<td>NR3C1</td>
<td>5q31.3</td>
<td>rs1048261</td>
<td>Serotoninergic, glucocorticoidergic</td>
<td>53</td>
</tr>
<tr>
<td>Dopamine receptor 3</td>
<td>DRD3</td>
<td>3q13.31</td>
<td>rs6280</td>
<td>Dopaminergic</td>
<td>60</td>
</tr>
<tr>
<td>Dopamine B hydroxylase</td>
<td>DBH</td>
<td>9q34</td>
<td>rs1611115</td>
<td>Dopaminergic</td>
<td>61</td>
</tr>
<tr>
<td>Brain-derived neurotropic factor</td>
<td>BDNF</td>
<td>11p14.1</td>
<td>rs6265</td>
<td>Dopaminergic</td>
<td>64</td>
</tr>
<tr>
<td>µ-Opioid receptor</td>
<td>OPRM1</td>
<td>6q25.2</td>
<td>rs1799971</td>
<td>Opioidergic</td>
<td>68</td>
</tr>
<tr>
<td>Fatty acid amide hydrolase</td>
<td>FAAH</td>
<td>1p33</td>
<td>rs324420</td>
<td>Endocannabinoidergic, opioidergic</td>
<td>72</td>
</tr>
<tr>
<td>5-Hydroxytryptamine transporter</td>
<td>SLC6A4</td>
<td>17q11.2</td>
<td>rs4251417</td>
<td>Serotoninergic</td>
<td>53</td>
</tr>
<tr>
<td>5-Hydroxytryptamine receptor 2A</td>
<td>HTR2A</td>
<td>13q14.2</td>
<td>rs2296972, rs622337</td>
<td>Serotoninergic</td>
<td>53</td>
</tr>
<tr>
<td>Tryptophan hydroxylase-2</td>
<td>TPH2</td>
<td>12q21.1</td>
<td>rs4570625</td>
<td>Serotoninergic</td>
<td>73</td>
</tr>
<tr>
<td>Serotonin transporter gene-linked polymorphic region</td>
<td>5-HTTLPR</td>
<td>17q11.2</td>
<td>Variable number tandem repeats (VNTR)</td>
<td>Serotoninergic</td>
<td>73</td>
</tr>
</tbody>
</table>

SNP, single nucleotide polymorphism.

Dopamine signalling pathways

Dopamine signalling pathways, which are dopaminergic projections, are the sets of projection neurons that synthesize and communicate the neurotransmitter dopamine. Dopamine neurotransmission is involved in pain modulation, mood regulation, and reward systems. Variants in genes that regulate dopamine metabolism and signalling have been linked to altered placebo responses.

**Monoamine oxidase (MAOA)**

MAOA is involved in the degradation of biogenic amines, including serotonin and dopamine. Variants in the MAOA gene have been associated with placebo responses and have been suggested to influence the placebo effect by modulating dopaminergic signalling.

**Catechol-O-methyltransferase (COMT)**

COMT is responsible for the degradation of catecholamines, including dopamine. A common genetic variant (rs4680) in the COMT gene, which changes valine to methionine at codon 158, has been linked to placebo responses. Individuals with the AA allele have been found to have reduced placebo responses compared to those with the GG allele. This variant is associated with decreased dopamine levels in the prefrontal cortex, which may influence placebo responses.

**Dopamine receptors**

Dopamine receptors are subdivided into D1-like (D1 and D5) and D2-like (D2-D4) subtypes. Variants in these genes have been linked to placebo responses, with individuals who have certain receptor subtypes showing reduced placebo responses.

**5-HTTLPR**

The serotonin transporter gene-linked polymorphic region (5-HTTLPR) is a polymorphism that influences the expression of the serotonin transporter gene. Variants in this region have been associated with placebo responses, with the VNTR (variable number tandem repeats) structure of the polymorphism showing varying effects on placebo responses.

Furthermore, the endocannabinoid pathway has been suggested to play a role in placebo responses, with genetic variants that influence the endocannabinoid system, such as Fatty Acid Amide Hydrolase (FAAH), also being associated with placebo responses. The implication of these genetic targets in the placebo effect is more limited.
study was based on a previous RCT with three groups, that is, an NTC group, a placebo acupuncture group and a placebo acupuncture+warmth caring group, to evaluate placebo acupuncture treatment of IBS. The results of that RCT showed that the best placebo treatment induced the strongest symptom relief since the RCT design potentially ruled out some factors affecting the responses to placebo treatment. Further, the genetic analysis results suggested that subjects in the homozygote of the rs4680 AA genotype with low enzyme activity resulting in high levels of dopamine in the body had the greatest placebo responses, and the G allele homozygous patients with high enzyme activity had the lowest placebo response. The heterozygotes of the GA genotype had an intermediate response. Moreover, another SNP rs4633, in the linkage with rs4680, had been found to give similar results.

To the best of our knowledge, to date the largest study of genetic variations in RCT patients with placebo and bupropion treatments for major depressive disorder examined a total of 532 variants in 34 candidate genes. Although there were no results for rs4680 in that study, several other SNPs in COMT were found to be significantly associated with placebo effects. However, after the correction for multiple comparisons, these SNPs were not found to be associated with placebo effects. Interestingly, in a recent laboratory study, the G allele of rs4680 related to the high enzyme activity of COMT was found to be significantly associated with a higher frequency of nocebo effects and complaint record. The finding suggests that when individuals with the GG genotype show absence of any significant improvements in symptoms, they may have more side effects, such as complaints and nocebo effects. However, this result must be verified in a wide population since the study primarily tested Caucasian women.

In addition to COMT, there are several other genetic target candidates of placebo effects in the dopamine pathway. The monoamine oxidase A (MAOA) gene encodes an enzyme bound to the outer membrane of mitochondria in most cell types and can catalyse the oxidative deamination of amines, such as dopamine, serotonin and norepinephrine, and thus takes part in reward pathways and affects serotonergic signalling pathways. A common SNP rs6323 (G>T) in MAOA has been found to reduce the enzymatic activity by 75% in individuals with the only allele of T. The first association study of rs6323 in the gene MAOA with placebo effects recruited patients with clinical depression from four placebo-controlled RCTs. Individuals containing the low-activity MAOA genotypes, which cause higher basal dopamine level, had higher placebo effects. However, in this study, rs4680 in the COMT gene was not found to be significantly associated with placebo effects, which may be because of a lack of statistical power, the subject difference or study design without a NTC group. In the above-mentioned largest genetic association study of placebo treatment, which sacrificed statistical power to detect variants with significant associations of placebo effects, the SNP rs6609257 within the gene MAOA involved in dopamine basal tone as well as rs1048261 in the nuclear receptor subfamily 3 group C member 1 (NR3C1) gene was found to be significantly associated with placebo-induced improvement in depression. The NR3C1 protein is a glucocorticoid receptor usually staying in the cytoplasm. On ligand binding, NR3C1 can be transported into the nucleus and functions both as a transcription factor and regulator of other transcription factors to mainly regulate the transcription of glucocorticoid responsive genes. These findings support that the MAOA and NR3C1 genes should be considered as molecular targets of placebo effects.

The dopamine receptor 3 (DRD3) encodes the D3 subtype of the five dopamine receptors, which are mediated by G proteins primarily located in the olfactory tubercle, nucleus accumbens and islands of Calleja in the brain, and are involved in cognitive, emotional and endocrine functions. A common exonic rs90280 (C>T) in the gene DRD3 can cause an AA substitution of glycine to serine at codon 9 (Gly9Ser). The mutant DRD3 with serine has been found to have a lower affinity for dopamine. A recent RCT of a novel drug (ABT-925) for treating schizophrenia examined the effects of variants in the DRD3. Patients with homozygous T allele of rs6280 in DRD3 gene were found to have significantly better outcomes in the group of placebo treatment than in the group of ABT-95 treatment with increasing doses. That study demonstrated that DRD3 should be a molecular target of the placebo effects, and supports that subjects homozygous for rs4680 A allele in the COMT gene have a greater placebo response.

The dopamine beta-hydroxylase (DBH) is an oxidoreductase belonging to the copper type II, ascorbate-dependent mono-oxygenase family. DBH is mainly expressed in neurosecretory vesicles and chromaffin granules of the adrenal medulla, and converts dopamine to norepinephrine. This enzyme has two forms, that is, soluble and membrane-bound forms. DBH has been reported to be associated with deficits in autonomic and cardiovascular function and psychiatric diseases. In the alcohol dependence studies, individuals with homozygous C allele of the rs1611115 in the DBH gene appeared to have better symptom improvement on the group of placebo treatment than on the group of naltrexone treatment. DBH was also examined in the largest genetic association study of the placebo and bupropion treatment for the major depressive disorder mentioned above. The SNP rs2873804 in the DBH gene was found to be significantly associated with placebo effects after the correction for multiple comparisons, which reinforces DBH as a potential molecular target for a placebo effect.

The brain-derived neurotrophic factor (BDNF) is a member of the nerve growth factor family of proteins. BDNF may promote neuronal survival and differentiation in the peripheral and central nervous systems, and participate in the modulation of axonal and dendritic growth.
and morphology. Being a major regulator of synaptic transmission and plasticity in the adult brain, BDNF has versatile roles in a range of adaptive neuronal responses, such as long-term depression, long-term potentiation, as well as homeostatic regulation of intrinsic neuronal excitability. Furthermore, BDNF has been found to regulate the stress response, modulate the pathogenesis of mood disorders and maintain turnover of dopamine. The functions of BDNF in neuroadaptive change and response to reward stimuli have attracted researchers’ interest in identifying plausible candidates involved in placebo effects. The SNP rs6265 (C→T) in the BDNF gene causes a valine to methionine substitution at codon 66 (Val66Met), which results in inefficient BDNF trafficking to secretory granules. Furthermore, the C allele of rs6265 has been found to be associated with greater placebo-induced activation of dopamine receptors D2 and D3 than that of T allele carriers; however, this gene was not found to be significantly associated with placebo analgesia despite that the SNP rs6265 has been hypothesised to reduce activity-dependent BDNF release.

Collectively, the results of association studies of placebo effects with dopamine-related genetic variants support the genes involved in dopamine pathway as molecular targets of placebo effects. More research with larger samples in studies that include NTCs would help provide definitive results.

**Opioid signalling pathways**

This system consists of multiple molecular signals generated by an opioid receptor binding to its physiological ligands. Opioids, broadly used as painkillers, are chemical substances extracted from opium (morphine, codeine, and so on), which possess strong analgesic and sedative effects. Opioid receptors are G-protein coupled receptors that are widely distributed throughout the human body and crucially involved in pain signalling in the central and peripheral nervous systems, and immunological response, and so on. There are four classes of receptors: mu (µ-opioid receptor, OPRM), kappa, delta and nociceptin. In studies of placebo analgesic effects, both the endogenous opioid and dopaminergic signalling pathways have been found to be activated. Especially, in antinociceptive responses to placebo, opioid receptor signalling has been found to be entangled with the dopamine signalling pathways.

Furthermore, OPRM1 has been found to affect the clinical outcomes of pain treatment in studies of placebo analgesia. The mutation of the A allele to G allele of rs1799971 in OPRM1 can cause an asparagine to aspartic acid alteration at codon 129 receptor, which may reduce the expression and function of OPRM1. The functional aspartic acid (G) allele of rs1799971 carriers has been found to have lower placebo-involved activation of dopamine neurotransmission unlike the asparagine (A) allele homozygotes, which suggests that genetic variation in OPRM1 could also contribute to variability of the placebo effects. In that same study, using positron emission tomography (PET) technology and radio tracers to label µ-opioid and dopamine receptors, compared with G allele carriers, the allele AA homozygotes of the functional rs1799971 showed an increase in the baseline level of OPRM in brain areas in response to pain and mood. Following a placebo treatment, G allele carriers were correlated with higher Neuroticism Extraversion Openness (NEO) personality scores and showed lower levels of mood, OPRM and dopamine receptor activation in the thalamus, nucleus accumbens and anterior insula. These findings implicate OPRM1 in the placebo-involved modulation and individual differences in neurotransmission. However, association studies of genetic variation in OPRM1 with the addictive effects of opioid drugs and psychostimulants (eg, amphetamine) have obtained conflicting outcomes. That conflict may provide some support for the need of an NTC to determine the genetic variation caused by differences in placebo effects.

**Endocannabinoid and serotonin signalling pathways**

Endocannabinoids (endogenous cannabinoids) are endogenous lipid-based retrograde neurotransmitters and include at least five derivatives of arachidonic acid, such as arachidonoyl ethanolamide and 2-arachidonoyl glycerol. They are released from postsynaptic neurons and bind to cannabinoid receptors, specifically cannabinoid type 1 (CB1) and CB2. The endocannabinoid signalling pathways are involved in regulating a variety of physiological and cognitive processes (such as appetite, pain sensation, mood and exercise-induced euphoria) and in mediating the pharmacological effects of cannabis. CB1 receptors are predominantly expressed in the peripheral and central nervous systems, and are mainly activated by the endocannabinoids, anandamide, as well as its mimic phytocannabinoid, tetrahydrocannabinol. Antagonist-based placebo analgesia studies have supported that endocannabinoid is involved in placebo analgesia.

The fatty acid amide hydrolase (FAAH) located on chromosome 1p33 is an integral membrane protein, which is responsible for the hydrolysis of some bioactive fatty acid amides, such as the neuromodulatory compounds anandamide and oleamide. Serving as the major degradative enzyme of endocannabinoids, FAAH may play roles in endocannabinoid responses to pain and placebo analgesia. The genetic variation in FAAH has been examined in a small study with the same subjects as mentioned above. The SNP rs324420 (C>A) within the FAAH gene encodes a missense substitution of proline to threonine at codon 129 (Pro129Thr). It has been reported that in response to pain subjects with homozygotes for the C allele of rs324420 had increased endocannabinoid levels in the brain, and a greater placebo analgesic response, and improved mood. These findings support that the endocannabinoid pathway genes are potential candidate molecular targets of placebo effects that are worth exploring further.

Serotonin is an important hormone and neurotransmitter with many roles. The serotonin signalling pathways...
are a set of projection neurons in the brain, including rostral and caudal groups, which synthesize and communicate the monoamine neurotransmitter serotonin. Individual neurons in these pathways are called serotonergic neurons. Since the serotonergic neurons innervate wide places, these pathways regulate mood, appetite and sleep and are relevant to many psychiatric and neurological disorders.

Since the high incidence of placebo effects in RCTs of placebo treatments for mood disease, the serotonin pathway could plausibly be examined for possible placebo effect-related genes. In the above-mentioned study, in which the association of 34 possible genes was examined with placebo effects, several genes involved in the serotonergic pathway were significantly related to placebo remission, including solute carrier family 6 member 4 (SLC6A4) rs4251417 and 5-hydroxytryptamine receptor 2A (HTR2A) rs2296972 and rs622337. SLC6A4 is an integral membrane protein that can terminate the action of serotonin and recycle it in a sodium-dependent manner through transporting the serotonin from synaptic spaces into presynaptic neurons. HTR2A is a guanine nucleotide-binding protein (G-proteins) coupled receptor for 5-hydroxytryptamine (serotonin) that plays a role in the regulation of behaviours, such as responses to anxiogenic situations and psychoactive substances, and in intestinal smooth muscle contraction and arterial vasoconstriction. Receptor ligand binding causes a conformational change in HTR2A that triggers intracellular signalling via G-proteins and modulates the activity of downstream effectors.

Moreover, serotonin-involved placebo effect genes have been examined in a social anxiety disorder (SAD) RCT. In the genetic PET study of SAD, a reduction in stress-related amygdala activity was accompanied by a reduction in anxiety symptoms under the treatment of placebo. Subjects homozygous for rs4570625 T allele within the tryptophan hydroxylase-2 (TPH2) gene promoter and the long allele of the serotonin transporter-linked polymorphic region (5-HTTLPR) have such a reduction. TPH2 is a member of the pterin-dependent aromatic acid hydroxylase family, which catalyses the first and rate-limiting step in the biosynthesis of serotonin. 5-HTTLPR is a polymorphic region located within the 5' regulatory region of the SLC6A4 gene including the promoter, which is composed of 16 tandemly repeated units in a long (L) allele, and deletion of repeat units 6–8 in a short (S) allele, where each repeat unit is 20–23 bps in length. The 5-HTTLPR region may have both positive and negative-acting cis-regulation on the expression of SLC6A4. Moreover, the SNPs of rs25531 and rs25532 have also been identified within 5-HTTLPR.

Although most genetic association studies of placebo effects have several limitations, such as small size and no NCT, the genes in Table 1 are potential molecular biological components of placebo effects in different disorders. However, more data are required to precisely define the roles of dopamine, opioid, endocannabinoid and serotonin based on the genetic background of placebo effects.

Recently, a placebo analysis based on the known seed genes influencing placebo effects tried to identify a subnetwork of interacting proteins involved in placebo effects. In this analysis, a placebo module constructed with an interactome of genes or proteins was identified to be significantly close to the vascular disease modules. Moreover, diseases with molecular network modules very close to the placebo module might be candidates for placebo as potential 'drugs'. For drugs targeting placebo module molecules, placebo effects may influence the drug test outcome. Furthermore, genetic variants in placebo module genes that modify the placebo effects, such as COMT, may provide a genetic target or biomarker of place effects. In this study, an indirect cohort was used to validate these findings by examining whether the placebo module had more genes with SNPs significantly related to a placebo effect than a random situation. Although a strict validation is required, this study suggests that the interaction between diseases/drug targets and the placebo module would tend to be stable. Given the complex interplay of expectation, behaviour and disease, a potentially complex network in the context of genes and environmental factors may determine placebo effects. Considering the complexity and lack of sufficient data, additional placebo studies are required.

The placebo effect is complex, and its physiology is incompletely understood. In previous studies, multiple intersecting pathways were found to be integrated into four signalling systems. In the viewpoint of Colloca and Miller, the core factors identified so far require further investigations to understand how they interact in complex biological networks.

CONCLUSION AND PERSPECTIVE

Placebo has a long history, but research on the genetics related to the placebo effect is in the early stage. In particular, research on placebome is in its infancy. A number of behavioural and neurobiological studies have explored the mechanism of placebo effects, including how they are stimulated. But the underlying mechanisms of placebo effect have not been fully understood. Although some genetic studies have been rigorously conducted, very few have explored the multiple possible genes in one time, like placebo studies. Then a primary limitation of this review is that it relies on a limited number of studies, so caution is required when considering these very preliminary conclusions. Although the molecules in the dopamine, opioid, serotonin and endocannabinoid systems might be targets for involvement in placebo effects on the basis of the combination of behavioural, neurobiological and genetic findings, placebo studies are required to identify genetic targets in the scope of the entire genome. Although bioinformatics analysis has identified the molecular placebo module, placebo studies with NTCs are also required to validate the related findings.

Contributors LC conceived and wrote the whole manuscript. LH proofread the manuscript.

Funding This work was supported by the Ministry of Science and Technology Precision Medicine Project (No 2017YFC0909200 and No 2017YFC1001300) and the Natural Science Foundation of Shanghai (No 19ZR1427700).

Competing interests None declared.

Patient consent for publication Not required.

Provenance and peer review Commissioned; externally peer reviewed.

Data availability statement Data is available from the authors on request

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

REFERENCES

Lei Cai is an associated professor of Shanghai Jiaotong University, information commissioner offiuatong University library, a youth editor for the Journal "Chinese medical ethics". He obtained the Ph.D degree from Fudan University, China and received Science trainings from Harvard Schooof public health, US. His research interests are focused on study the molecular basis of mind-body interaction and Big data analysis. Till now, over 40 papers and four chapters for two books have been published, two invention patents have been authorized. He has been PI of several projects or research tasks, such as National Natural Science Foundation of China/ Shanghai, the Ministry of Education (MOE) Scientific Research Foundation, and the key research and development program of the Ministry of Science and Technology.