Implications of Placebo and Nocebo Effects for Clinical Practice: Expert Consensus

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Placebo effect · Nocebo effect · Patient’s expectancies · Clinical practice · Patient-clinician communication · Evidence-based ethical recommendations

Abstract
Background: Placebo and nocebo effects occur in clinical or laboratory medical contexts after administration of an inert treatment or as part of active treatments and are due...
Implications of Placebo and Nocebo Effects for Clinical Practice

Introduction

Placebo and nocebo research has now evolved from a methodological consideration within clinical research to a distinct and expanding interdisciplinary field in its own right. Placebo studies encompass a broad variety of disciplines from biomedicine and neuroscience to the social and behavioural sciences and the humanities [1–14]. Placebo and nocebo effects refer to the beneficial or adverse effects that occur in clinical or laboratory medical contexts, respectively, after administration of an inert treatment or as part of active treatments, due to mechanisms such as expectancies of the patient. Current research incorporates both the study of placebo and nocebo responses as an outcome in clinical trials as well as the study of placebo and nocebo effects and their psychological and neurobiological mechanisms in various clinical applications, including many medical disciplines. Current objectives in the field are to produce cross-disciplinary conceptualizations of placebo and nocebo effects; to re-evaluate the ethics of placebos in clinical trials and practice; and to initiate the ethical translation of empirical findings into clinical practice.

Robust empirical evidence now demonstrates that placebo and nocebo effects are both significant and measurable for many conditions (e.g., pain, depression, Parkinson’s disease, fatigue, allergies, and immune deficiencies) [1]. Importantly, placebo and nocebo effects can substantially modulate the efficacy and tolerability of active pharmacological or other medical treatments [15, 16]. This empirical evidence from experimental and clinical studies challenges health-care professionals to translate and implement the findings of placebo and nocebo research into practice. This translation relates not only to the possible use of placebos as part of regular treatments (e.g., placebo pills) but, far more importantly, to the systematic utilization of the mechanisms underlying placebo and nocebo effects to augment established treatment strategies (e.g., attention to expectations and empathy in patient-clinician communication).

Due to the interdisciplinary nature of this emerging field with its potentially wide applications across health care, there is a need to develop and formalize guidelines on the available evidence, including the possible implications of placebo and nocebo effects for clinical practice [17]. While the American Medical Association (AMA) provides explicit guidelines on the clinical use of placebos, the General Medical Council (GMC) in the UK provides no such ethical guidelines [18]. Notwithstanding the content of ethical guidelines and codes, studies reveal that the use of both pure (e.g., sugar pill) and impure (e.g., antibiotics for viral infections) placebos by doctors is widespread [19–21]: for example, in the US, 55% of internists and rheumatologists reported using placebos; in the UK, 77% of primary care doctors reported that they used placebos at least once per week, while 86% of primary care doctors in Denmark admitted that they had used placebos at least once within the last year. Surveys aimed at investigating patients’ attitudes reveal that the majority of respondents believe that placebo use by doctors is acceptable under certain circumstances [22]. Until now, however, there has been no study of expert opinions in placebo and nocebo studies about the implications of this research for clinical practice.

To address this gap, a clinical expert meeting was organized to survey established placebo researchers’ views about placebo and nocebo effects and their translation...
into clinical practice. The aim of this survey and meeting was to develop a consensus on clinical recommendations based on the current state of the art in placebo and nocebo research among experts in the field.

**Methods**

**Expert Group**

Of the 31 invited speakers at the 1st official Society for Interdisciplinary Placebo Studies (SIPS) conference, 29 agreed to take part; the 2 who chose not to participate indicated that their research was not clinically focused. The 29 invited experts completed the survey, of whom 4 invited speakers were unavailable to participate due to personal circumstances. A workgroup of 6 members (A.W.M.E., L.C., C.B.; R.K., T.J.K., and J.M.K.) prepared the survey and expert meeting.

The 29 participants were from 12 different countries, had an average age of 49.1 years (SD 11.1), and 45% were female. In total, 65.5% worked clinically (17.2% physicians, 41.4% psychologists, and 6.9% acupuncturists). The participants’ backgrounds included anaesthesiology, neurology, cognitive neuroscience, primary care, internal medicine, psychiatry, psychosomatic medicine, health and medical psychology, clinical psychology, epidemiology, medical ethics and philosophy. The participants had on average 14.2 years (SD 8.0) of research experience in the field of placebo and nocebo studies or related areas since their doctoral degree.

**Survey**

Based on a literature review of the relevant empirical evidence on possible clinical applications of placebo and nocebo effects, a survey was developed that focused on 4 themes: (a) prescription of placebo as regular treatment; (b) open-label prescription of placebo; (c) nocebo effects, and (d) patient-clinician communication. Subsequently, items for each theme were developed and checked by all members of the workgroup for relevance, readability and clarity. This process yielded 40 items in all, 10 for each theme. Participants were asked to read each statement and rank it on a 0- to 10-point scale with 0 meaning “totally disagree” and 10 meaning “totally agree.” In addition, participants had the opportunity to provide written comments on each item (see online suppl. Appendix Table S1 for a complete overview of the survey; for all online suppl. Material, see www.karger.com/doi/10.1159/000490354). The survey was analysed by calculating the mean score for each item. High agreement with a statement was defined as a mean score ≥8, and high disagreement with a statement was defined as a mean score ≤2 on the 10-point scale. Items that resulted in more mixed levels of agreement (scores between 2 and 8) were not discussed at the clinical expert meeting.

**Clinical Expert Meeting**

The clinical experts met in a 1-day pre-conference session in Leiden, The Netherlands, on April 2, 2017. At this meeting, the results of the survey and possible recommendations and discussion points were explored in plenary sessions. Consensus was reached based upon results of the survey and the discussion during the meeting. The meeting was audio-recorded and minutes were taken.

<table>
<thead>
<tr>
<th>Table 1. Summary of the recommendations formulated by the expert group</th>
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<tr>
<td><strong>Dos</strong></td>
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<tr>
<td>1. Consider placebo effects as part of regular treatment</td>
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<tr>
<td>2. Inform patients about placebo and nocebo effects in such a way that treatment effects are maximized and side effects are minimized</td>
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<tr>
<td>3. Ensure a patient-clinician relationship that is characterized by trust, warmth and empathy in order to maximize placebo effects and minimize nocebo effects</td>
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<td>4. Train health-care providers in patient-clinician communication to maximize placebo effects and minimize nocebo effects</td>
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<td>5. Prefer open-label rather than hidden placebo prescription in those cases where there is evidence for efficacy and where prescribing a placebo is legal</td>
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<td><strong>Don’ts</strong></td>
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<tr>
<td>1. Do not take risks (e.g., prescribing invasive treatments) to maximize placebo effects</td>
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<td>2. Do not consider deception a necessary component of placebo effects</td>
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**Results**

In the sections below, we describe the main results of the survey and expert meeting. The initial discussion centered on defining important components of the placebo concept. We then summarize the survey statements with high agreement, as well as the related evidence, together with relevant comments discussed at the clinical expert meeting. A summary of the recommendations is listed in Table 1.

**Conceptualization of Placebo Effect and Placebo Response**

The experts agreed that it is crucial to distinguish between placebo and nocebo responses versus placebo and nocebo effects. In line with the drug terminology proposed by Fisher et al. [23] (1965), the placebo and nocebo response includes all health changes that result after administration of an inactive treatment (i.e., differences in symptoms before and after treatment), thus including natural history and regression to the mean. The placebo and nocebo effect refers to the changes specifically attributable to placebo and nocebo mechanisms, including the neurobiological and psychological mechanisms of expectancies. These mechanisms are shaped, for example, by
verbal instruction, or nonverbal or situational cues that affect treatment expectancies (see communication by the SIPS community, July 29, 2016) [see also 24]. Importantly, placebos and nocebos not only have effects during the prescription of placebo pills, but they can also substantially modulate the efficacy and tolerability of active pharmacological or other medical treatments. There was strong consensus that recommendations should be based on the research evidence on placebo and nocebo effects, including the involved mechanisms (e.g., expectancies) and their consequences for medical practice (e.g., patient-clinician communication).

Results of the Survey and Expert Meeting
Considering Placebo Effects as Part of Regular Treatment

There was a consensus regarding the importance of making optimal use of placebo effects to achieve better treatment outcomes (A1) and of optimally informing patients about placebo effects (A8), for example, to explain that a patient might improve due to factors other than the treatment itself, such as expectancies regarding treatment prognosis. The consensus was based primarily on the broad evidence that now exists for placebo and nocebo effects on clinical, neurobiological and laboratory outcomes, at least for the areas of pain, depression, Parkinson’s disease, fatigue, allergies and immune deficiency [1–14]. However, the experts also acknowledged the current lack of knowledge about how best to provide this information to patients to ensure that patients are optimally informed. For example, there is insufficient knowledge about the ways in which patients and health-care providers can maximize placebo effects and minimize nocebo effects over time during repeated interactions and treatments [25].

The experts strongly agreed that clinicians should not prescribe or practice more invasive treatments simply to engender more potent placebo effects (A7). Although some systematic reviews have found that more intensive and invasive treatments can augment placebo effects, results are not consistent and are often moderated by factors such as the symptom or condition treated or the specific procedures used [26]. Moreover, there are clear practical and ethical restrictions on prescribing more invasive treatments for the sake of producing stronger placebo effects, since these procedures are often more expensive and entail a higher risk of undesirable side effects. Consequently, the experts agreed that prescription of more intensive and invasive treatment (e.g., injections instead of pills) to increase placebo effects is not justified.

Open-Label Placebo

Where open-label placebo (i.e., prescription of placebo pills with the knowledge and assent of the patient) was concerned, there was relatively high consensus that these should be preferred to hidden placebos (B3). Importantly, there are compelling ethical arguments for informing patients about placebo treatments in clinical contexts [27]. Related to these findings, the majority of the experts strongly agreed that deception is not necessary for placebo effects to occur (B2). This viewpoint is empirically supported by recent studies in patients with irritable bowel syndrome and chronic pain, which showed beneficial effects of open-label placebo on primary symptom outcomes, such as pain or disability [28–31]. However, more research is needed in other conditions and patient groups before recommendations can be developed about their possible use in clinical practice. For example, non-significant findings were found in a pilot study in patients with a major depressive disorder in a small sample of 12 patients, although the medium effect size was comparable to the other trials [32–34]. It has to be established whether specific patient groups or subgroups of patients benefit more than others from open-label placebos, or whether biological or psychological markers are associated with the effectiveness of these open-label placebo strategies. Promising is the case of dose-extending placebos (e.g., placebo given along with active medication) that mimic the psychobiological responses that are associated with the effectiveness of the medications with the potential of reducing side effects and costs [32]. Although prescription of placebos is not yet regulated in any part of the world and additional research is necessary on its short- and long-term effects, open-label prescription of placebos could be considered as a possible future watch and wait strategy or for use in long-term conditions when other treatment options have failed [33].

Nocebo Effects

There was a consensus regarding strategies to minimize and prevent nocebo effects. Experts agreed that nocebo effects should be explained to patients (C1) and that information about side effects should be presented in such a way that nocebo effects are minimized (C4). The relatively strong consensus regarding optimally informing patients to minimize nocebo effects might be due to its high clinical relevance, which is based on findings that nocebo effects have been shown to consistently worsen treatment outcomes and to be at least in part responsible for side effects [6, 10–12, 35]. Indeed, there is convincing evidence that the way in which patients are informed
about risks and side effects influences the likelihood of their occurrence. For example, when reassuring words are used during a local anaesthetic injection instead of emphasizing the pain experience during the procedure, pain reports can be reduced [36]. However, there are several ethical considerations that are required for optimal informed consent of patients, including balancing the need for honesty and transparency with the requirement that harms should not be induced or increased unnecessarily. For example, tailored strategies might be applied after the necessary information has been offered, depending on the patient’s need for more detailed information on the risks and side effects of treatment. Such strategies might be particularly useful for those patients who have a high risk of developing nocebo effects, such as individuals who exhibit high levels of somatic amplification or fear of side effects [7, 10, 11, 35]. Moreover, evidence is needed about the consequences of systematically informing patients about the role of nocebo effects, for example, by introducing the terminology of nocebo effects, or determining the optimal moment for informing patients about nocebo effects during ongoing treatments [37].

There was also consensus that clinicians should receive training and education to minimize nocebo effects (C8 and C9). Training and education might include informing health professionals about the negative impact of nocebo effects, as well as training them in strategies for optimal verbal and non-verbal communication, including information about risks and side effects [7, 9–11, 35–37]. Again, however, there was acknowledgement that there has been a lack of research into the education and training of health professionals in relation to nocebo effects. Future studies might focus on evaluating different types of educational strategies to obtain insight into the most effective tools for training professionals to minimize nocebo effects (e.g., how information is communicated) as well as into optimal modes of education and training (e.g., including virtual reality tools).

Patient-Clinician Communication

The results of the survey showed a high level of agreement regarding patient-clinician communication (4 of 10 survey items). Experts agreed that a good patient-clinician relationship is essential to make optimal use of placebo effects to increase therapeutic efficacy (D2). Indeed, evidence is mounting that characteristics such as trust, warmth, and empathy are helpful in medical communication to foster placebo effects [38, 39]. In addition, there was clear consensus that clinicians should receive regular education and training about how to make optimal use of placebo effects in their treatments (D6 and D7). However, studies examining which elements this training might include are scarce [17]. For example, a critical point in the patient-clinician communication is how to optimize expectancies without risking subsequent violation of patient’s expectations and how to avoid eliciting overly positive expectations that might harm trust in the treatment. An important step will be to study specific features of disclosures (e.g., content, setting) with the aim of ethically tailoring education and training towards maximizing placebo and minimizing nocebo effects. Finally, experts agreed that medical ethics education encompassing placebo and nocebo effects should be a routine part of clinician training (D8).

**Discussion**

Intensive research in recent decades provides substantial evidence for the potential benefits of placebo effects, as well as the possible harms of nocebo effects [1–14]. In this survey of international experts on placebo research, there was strong consensus that patients should be informed about placebo and nocebo effects and that healthcare professionals should be trained to maximize placebo effects and minimize nocebo effects. The current paper forms a first step towards developing evidence-based and ethical recommendations about the implications of placebo and nocebo research for medical practice, based on the current state of evidence and the consensus of experts.

More research is needed in areas such as the prescription of open-label placebos, the optimization of intervention strategies to maximize placebo effects and minimize nocebo effects, and generalizability across different conditions. For example, it is important to ascertain which specific strategies for informing and educating patients and health professionals work best, as well as the optimal timing for the delivery of these interventions. It is also relevant to study the possible additive or interactive effects of different ways to maximize placebo effects, such as optimally informing patients, changing environmental cues or using conditioning during medical procedures. This might be particularly urgent in the area of nocebo effects, for example, for those patients who have a higher risk of developing severe adverse events due to high levels of fear of side effects or previous traumatic medical procedures. Large-scale research should also be encouraged with the aim to understand the substantial individual differences between placebo and nocebo responders and the possible neurobiological, psychological and genetic pre-
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