Use of homeopathic preparations in experimental studies with healthy plants

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Background: The last comprehensive review of experimental research on effects of homeopathic treatments on plants was published in 1984, and lacked formal predefined criteria to assess study quality. Since then several new studies with more advanced methods have been published.

Objectives: To compile a review of the literature on basic research in homeopathy with healthy plants with particular reference to studies investigating specific effects of homeopathic remedies.

Methods: The literature search included English, French, German, Italian, Portuguese and Spanish publications from 1920 to April 2009, using predefined selection criteria. We included experiments with healthy whole plants, seeds, plant parts and cells. The outcomes had to be measured by established procedures and statistically evaluated. We developed a Manuscript Information Score (MIS) and included only publications which provided enough information for proper interpretation (MIS ≥ 5). A formalised Study Methods Evaluation Procedure (SMEP) was used to evaluate these studies, and the subgroup of studies with adequate controls to identify specific effects.

Results: A total of 86 studies in 79 publications was identified, 43 studies included statistics, 29 had MIS ≥ 5, and 15 studies investigated the specificity of homeopathic preparations. Specific effects of decimal, centesimal and fifty millesimal potencies were found including dilution levels far beyond the Avogadro number. In consecutive series of potencies only some of the tested potencies showed effects. There were many individual studies with diverse methods and very few reproduction trials.

Conclusions: Healthy plant models seem an useful approach to investigate basic research questions about the specificity of homeopathic preparations. More investigations with more advanced methods are recommended, especially in the sectors of potentisation techniques, effective potency levels and conditions for reproducibility. Systematic negative control experiments should become a routine procedure to control the stability of the experimental systems. Homeopathy (2009) 98, 228–243.

Keywords: Review; Basic research; Homeopathy; Potentisation; Potentised dilutions

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Introduction

The last comprehensive review of experimental research on the effects of homeopathic treatments on plants was authored by Scofield in 1984. Since then several new investigations have been published, and more advanced methods used. In addition, the review by Scofield did not use any formal predefined criteria to assess study quality such as the criteria introduced by Becker-Witt for physico-chemical research and Bluth and Witt for in vitro research in homeopathy.

It is thus interesting to compile a new review of the literature about the effects of homeopathic treatments of plants. This would make it possible to determine the current state of research and to base future studies on research questions raised by results of prior work. We used predefined criteria to include only the publications which provided enough information to be interpreted properly. Furthermore, we focused on identifying studies which investigated specific effects of homeopathic remedies, i.e. effects related to the substance potentised.

Homeopathic basic research with plants can be divided into three major fields: experimental models with healthy plants, poisoned or impaired plants, and infected plants (phytopathological models). This publication will review the studies which experimented with healthy plants. The review about phytopathological models is published by Betti et al. in this issue, the review about studies with impaired plants will be published elsewhere.

Methods

Sources for the literature search

We retrieved most of the literature from the authors’ personal libraries. These literature collections were compiled by searching and collecting basic research articles for years, mostly by checking bibliographies of reviews and articles, by manually scanning scientific journals, and by information from colleagues. Additionally, the HomBRex Database (maintained by the Karl und Veronica Carstens-Stiftung, Essen, Germany) was used. Searching with the help of standard online literature databases (e.g. MEDLINE) was not very successful, because most studies are not indexed there.

Literature selection

This review covers publications that reported on experiments with homeopathy in healthy plants. This includes experiments with whole plants, parts of plants, plant cells and plant seeds. Studies were excluded if they featured with plants which were poisoned, infected, or stressed on purpose, e.g. by special experimental conditions such as the absence of light for seeds that require light for germination. Outcome parameters had to be measured by established procedures, e.g. length, weight, leaf area or secondary metabolites. Studies using unconventional methods such as Kirlian photography or bio photon emission as measurement techniques were not included in this review. Publications in German, English, French, Italian, Spanish or Portuguese were reviewed. In order to gather a comprehensive literature collection, we included all relevant publications from January 1920 to April 2009. Any earlier and later publications were excluded.

In some publications, multiple experiments or studies with differences in methods, set-up or results were described in one paper. If they were described separately, we decided to subdivide the publications into ‘studies’.

The studies identified by this selection procedure showed marked differences in methodology and manuscript information content. To compile a comparable and informative publication pool, we developed three further selection criteria.

Statistics: We excluded all publications which did not use a statistical evaluation of the results (minimum: mean/median, number n and standard deviation or standard error).

Manuscript information score (MIS): The MIS was developed to include only publications with sufficient information to be interpreted properly (see Table 1). In the MIS, a maximum of 10 points were given for five category groups. A minimum of 5 points was necessary for the study to be included in the review. All publications were independently evaluated by two reviewers. Any differences in rating were resolved by discussion.

Study Methods Evaluation Procedure (SMEP): The SMEP evaluates important features of the study set-up. Our focus was on the control samples, because adequate controls are of particular importance investigating specific effects of homeopathic preparations. We distinguished eight different types of control. In addition, four methodological categories (blinding, randomisation, number of independent experiments, systematic negative control experiments) were reviewed.

Studies that provide evidence of specific effects of homeopathic remedies, i.e. effects related to the mother tincture diluted and therefore implying some sort of ‘memory’ of the potentisation medium, have to be well designed to distinguish results from artefacts. Some authors suggested that silicates, other molecules and various ions are dissolved from the potentisation vessel during the succussion process. Unsuccessful potentisation medium contains fewer of these contaminants. Using an unsuccessful potentisation medium as sole control may generate false-positive results when it comes to identifying treatment effects that are specific to homeopathic remedies.

Essential controls to avoid these kinds of artefacts consist of working with a succussed potentisation medium or with a potentised potentisation medium (diluted and succussed in the same way as the potentised test substances). If the test substance first has to be dissolved or triturated in a medium other than the final potentisation medium, the most adequate control to identify specific effects is potentised solvent (e.g. potentised lactose or acetone without any other primary substance), provided that the solvent is potentised in the final potentisation medium. Further valid controls for identifying specific effects are other potentised test substances, if they follow analogous methods of production.

In addition to the controls just discussed, four further controls are itemised in the SMEP to depict the complete
list of controls chosen in the studies. The unsuccesful poten-
tisation medium is a common control in basic research
in homeopathy. In combination with a succussed or with
a potentised potentisation medium, it makes it possible
to investigate if there is a succussion effect on the potentisa-
tion medium. Dilutions of the test substance (diluted in
the same dilution steps as for potentisation but without suc-
cussion) can be regarded as a control to identify the impor-
tance of succussion in the potentisation process. In the
diluting process, however, properly stirring the dilution is
necessary to yield a homogeneous distribution of the mole-
cules in the dilution medium. It seems difficult to us to cre-
ate a precise distinction between stirring and potentisation.
The positive control is a material dose of a test substance
with a well-known effect. In comparison to the potencies,
it can be used to investigate the modification of the specific
substantial effect through potentisation. Few studies with
adult plants used a control group with no treatment.

Some older studies compared different potency levels of
the same test substance, without factoring another control
into the analysis.

We checked four further methodological key factors. We
determined whether the described experiments were carried
out under blind conditions. We also evaluated whether arte-
facts that might be produced by differing local growing
conditions like temperature and light were avoided by
randomising the samples in time and space (e.g. pots with
plants or the germination vessels). Additionally we checked
whether the stability of the chosen experimental set-up was
demonstrated by systematic negative control experiments.
These are experiments with the same set-up as potency exper-
iments using only one control substance for all samples (e.g.
distilled water). With successfully standardised laboratory
conditions and homogeneous quality of seeds or plants, no
significant differences between the samples may be observed.

We also assessed the number of independent experiments:
does the study consist of only one experiment, or were there
multiple independent experiments being carried out?

All publications were independently evaluated by two
reviewers. Any differences in assessment were resolved
by discussion.

*Extraction of information:* The final step was thoroughly
extracting the information from the study reports. At this
point, studies that investigated specific effects were finally
selected and sorted.

## Results

In some publications, multiple experiments or studies
with differences in methods, set-up or results were
described in one paper. If they were described separately,
we decided to subdivide the publications into ‘studies’. A
total of 79 publications, including 86 studies, were identi-
fied. The first publication was by Kolisko in 1923,
whilst the newest by Scherr in 2009. Of these 79 publica-
tions (86 studies), 43 either did not use a statistical analysis
to evaluate the results, or did not mention the statistics in the
publication. 36 publications (43 studies) with statistics
(published from 1962 to 2009) remained within the review-
ning procedure.

In the following we will always refer to the number of
studies (instead of the number of publications). We thus
have 43 studies from 36 papers which had some sort of
statistical evaluation and which were included in the further
reviewing process (for an overview, see Figure 1).

### MIS

29 of 43 studies achieved 5 or more points in the MIS.
They thus contained sufficient information for more de-
tailed interpretation. Table 2 gives an overview of the

### Table 1

<table>
<thead>
<tr>
<th>MIS</th>
<th>Fully described</th>
<th>Party described</th>
<th>Not mentioned</th>
</tr>
</thead>
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<tr>
<td>Score</td>
<td>2 points</td>
<td>1 point</td>
<td>0 points</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Experimental setup</th>
<th>Detailed information is given: way of treatment of plants, growth period, time of measurements, etc.</th>
<th>Only some details are described or few information about the set-up is given</th>
<th>No information is given about the experimental set-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Materials</td>
<td>All materials used in the experiments are described with trade name, etc.</td>
<td>Some materials used in the experiments are described or mentioned</td>
<td>No information is given about the materials used</td>
</tr>
<tr>
<td>Measuring instruments</td>
<td>Measuring instruments are described in detail, operation mode, trade name, type, etc.</td>
<td>Measuring instruments are only mentioned</td>
<td>There is no information about measuring instruments in the paper</td>
</tr>
<tr>
<td>Potentisation</td>
<td>Potentisation technique, date and time of potentisation and potentisation medium are described in detail</td>
<td>Some information about potentisation technique is given</td>
<td>No information about potentisation, only the potentised test substance is mentioned</td>
</tr>
<tr>
<td>Controls</td>
<td>Detailed information eg: sterile distilled water from the same batch of distilled water...</td>
<td>Some information about the sort of control is given: e.g.: water control</td>
<td>Controls are not mentioned or not done</td>
</tr>
</tbody>
</table>
studies, methods used, plant models, tested substances and potency levels, controls, and results the authors reported.

**SMEP**

By applying the SMEP to the 29 studies with MIS ≥ 5 we identified the controls applied as well as the following key methodological factors: blinding, randomisation number of independent experiments and systematic negative control experiments. Due to the heterogeneity of the items checked, rating of the studies with a quantitative sum score did not seem adequate to us. Depending on the objectives and set-ups of the studies, different preparations may be regarded as adequate controls. Our main focus was to identify the studies which used adequate controls to investigate specific effects of homeopathic preparations.

**Controls and specific effects:** Eighteen studies used only one control, while 11 studies included two or three controls. Of the 18 studies with one control, twelve studies used unsuccussed potentisation medium as the control and six studies one of the other types of control. Three of these 12 studies mentioned unsuccussed potentisation medium as control, but evaluated the results by comparing with other potency levels of the same test substance (see Table 2, column controls). Fifteen of the 29 studies included adequate controls to identify specific effects of the tested homeopathic remedies (see discussion below).

**Blinding and randomisation:** In the group of 29 studies with five or more points in the MIS, 11 studies were carried out under blind and randomised conditions. Four studies were blinded only. In six studies, the samples of the experiments were distributed at random, but the researchers were not blinded to the treatments. Eight of the 29 studies do not mention blinding or randomisation (see Table 2).

**Systematic negative control experiments:** Five studies described systematic negative control experiments to control any disruptive influences of inhomogeneous laboratory conditions (see Table 2, column Methods).

**Number of independent experiments:** Ten of 29 studies comprised only one experiment. Fourteen studies carried out two to eleven experiments included. Three studies consisted of 20–40 experiments. In two studies the number of independent experiments was not clearly identifiable in the manuscript (see Table 2).

**Combination of quality criteria for studies of specific effects:** Four of 15 studies with controls to identify specific effects used all quality criteria (blinding, randomisation, systematic negative control experiments, several independent experiments).55,56,85

**Extraction of information**

This part includes only the 29 studies with sufficient information available (MIS ≥ 5), including the studies, which investigated the specificity of homeopathic preparations (Table 2).

**Plant models:** The most frequently used experimental plant model was the seedling model. The influence of homeopathic potencies on the germination and the growth of the seedlings was investigated in 13 studies with wheat seeds,53,54,57,60,73,75–78 in three studies with dwarf peas,55,56 in one study with Sida rhombifolia67 and in one study with mung seed.62 A second experimental plant model investigated the influence of homeopathic potencies
<table>
<thead>
<tr>
<th>Author &amp; Year</th>
<th>Reference</th>
<th>Plant</th>
<th>Methods</th>
<th>Independent experiments</th>
<th>Tested substances</th>
<th>Tested potency levels</th>
<th>Controls</th>
<th>Effective potency levels–control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scherr 2009</td>
<td>85</td>
<td>Duckweed (Lemna gibba L.)</td>
<td>b; r; s</td>
<td>Gibberellic acid, Kinetin, Lemna m.; 5; Silver n.: 1</td>
<td>Gibberellic acid, Kinetine, Silver nitrate, Lemna minor</td>
<td>14x–30x</td>
<td>U + S</td>
<td>Gibberellic acid: 15x, 17x, 18x, 23x, 24x</td>
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<td>Sukul 2009</td>
<td>87</td>
<td>Lady's finger (Abelmoschus excelentus L. Moench)</td>
<td>b; r; s</td>
<td>CCC, CCC (nano), MH</td>
<td></td>
<td>30c, 200c</td>
<td>P</td>
<td>30c, 200c</td>
</tr>
<tr>
<td>Baumgartner 2008</td>
<td>56</td>
<td>Dwarf peas (Pisum sativum L.), cv. 'Früher Zwerg', harvests 1997–2000 Sida rhombifolia</td>
<td>b; r; s</td>
<td>8</td>
<td>Gibberellic acid</td>
<td>17x, 18x</td>
<td>U + S</td>
<td>17x</td>
</tr>
<tr>
<td>Marques 2008</td>
<td>67</td>
<td>Sida rhombifolia</td>
<td>r</td>
<td>5</td>
<td>Cymbopogon winterianus Jowitt (Citronella)</td>
<td>3c, 6c, 12c, 24c, 30c</td>
<td>U</td>
<td>3c, 6c, 12c, 24c, 30c</td>
</tr>
<tr>
<td>Sukul 2008</td>
<td>86</td>
<td>Pigeon pea (Cajanus cajan L. Millsp.)</td>
<td>b; r; s</td>
<td>CCC, CCC (nano), MH</td>
<td></td>
<td>30c, 200c</td>
<td>P</td>
<td>30c, 200c</td>
</tr>
<tr>
<td>Scherr 2007</td>
<td>84</td>
<td>Duckweed (Lemna gibba L.)</td>
<td>b; r; s</td>
<td>1</td>
<td>Silver nitrate, Copper sulphate, Gibberellic acid, Auxin, Kinetin, Lactose, Lemna minor, Methyl jasmonate, Metoxuron, Phosphorus, Potassium nitrate, Sulphur</td>
<td>14x–30x</td>
<td>U + S</td>
<td>Silver nit: 24x, 28x, 29x; Kinetin: 14x, 16x, 20x, 23x, 26x, 27x, 30x; Phos: 21x, 25x, 29x</td>
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<tr>
<td>Rossi 2006</td>
<td>79</td>
<td>Lettuce (Lactuca sativa), cv. 'Veronica'</td>
<td>r</td>
<td>1</td>
<td>Carbo vegetable</td>
<td>6c, 12c, 30c, 100c, 200c</td>
<td>U, (N)</td>
<td>6c, 12c, 30c, 200c</td>
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<tr>
<td>Baumgartner 2004-I</td>
<td>55</td>
<td>Dwarf peas (Pisum sativum L.), cv. 'Früher Zwerg'</td>
<td>b; r; s</td>
<td>Series 1: 4</td>
<td>Gibberellic acid, Kinetin, Auxin, Abscisic acid</td>
<td>I: 12x–30x</td>
<td>U + S; (P-C)</td>
<td>Gibberellic acid: 13x, 15x, 17x, 23x; Kinetin: 19x</td>
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<tr>
<td>Baumgartner 2004-II</td>
<td>55</td>
<td>Dwarf peas (Pisum sativum L.), cv. 'Früher Zwerg'</td>
<td>b; r; s</td>
<td>Series 2: 6; Series 3: 4</td>
<td>Gibberellic acid</td>
<td>II: 17x; III: 17x</td>
<td>U + S</td>
<td>17x</td>
</tr>
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<td>Chapman 2004</td>
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<td>Lettuce (Lactuca sativa), cv. 'Tom Thumb'</td>
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<td>Sulphur, Silicea</td>
<td>LM1</td>
<td>S</td>
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<td>Year</td>
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<td>Species</td>
<td>Potency Levels</td>
<td>Methods</td>
<td>Controls</td>
<td>Treatment Effects</td>
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<tr>
<td>2001</td>
<td>Andrade</td>
<td>Justicia pectoralis L. (Jacqu.)</td>
<td>b; r 1</td>
<td>Justicia; Acanthaceae; Cumarina P.A.; Guaco; Phos; Sulphur; Amica montana; Humic acid</td>
<td>3c</td>
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<tr>
<td>2000</td>
<td>Brizzi</td>
<td>Wheat (Tritium durum L.), cv. 'MEC'</td>
<td>b; r unclear</td>
<td>Arsenicum album</td>
<td>23x, 27x, 25x, 30x, 35x, 40x, 42x, 45x U; (D; P) 27x, 25x, 30x, 35x, 40x, 42x, 45x</td>
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<tr>
<td>1994-I</td>
<td>Betti</td>
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<td>b; r 10</td>
<td>Arsenicum album</td>
<td>23x, 30x U No effects</td>
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<td>1994-II</td>
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<td>Wheat (Tritium durum L.), cv. 'MEC'</td>
<td>b; r 8</td>
<td>Arsenicum album</td>
<td>23x, 25x, 30x, 35x, 40x, 45x (U; D) P 40x + 45x</td>
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<tr>
<td>1994a-I</td>
<td>Pongratz</td>
<td>Wheat, cv. 'Mephisto'</td>
<td>b unclear</td>
<td>Silver nitrate</td>
<td>24x–26x U 24x, 26x</td>
<td></td>
<td></td>
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<td>1994a-II</td>
<td>Pongratz</td>
<td>Wheat, cv. 'Mephisto'</td>
<td>b 2</td>
<td>Silver nitrate</td>
<td>24x P 24x</td>
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<td>1994a-III</td>
<td>Pongratz</td>
<td>Wheat, cv. 'Mephisto'</td>
<td>b 1</td>
<td>Silver nitrate</td>
<td>24x–26x U 24x, 26x</td>
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<td>1991</td>
<td>Endler</td>
<td>African violet (Saintpaulia sp.)</td>
<td>b; r 3</td>
<td>Indole butric acid</td>
<td>33x S 33x</td>
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<td>1990</td>
<td>Pongratz</td>
<td>Wheat, cv. 'Mephisto'</td>
<td>b 10</td>
<td>Silver nitrate</td>
<td>24x P 24x</td>
<td></td>
<td></td>
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<tr>
<td>1986</td>
<td>Chou</td>
<td>Mung seed</td>
<td>1</td>
<td>Fertilizer</td>
<td>1x–24x U; D; (P-C) Potency levels not mentioned 5c, 7c, 9c</td>
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<td>1979b</td>
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<td>1971</td>
<td>Pelikan</td>
<td>Wheat, own cultivation</td>
<td>r 40</td>
<td>Silver nitrate</td>
<td>8x–19x U 13x, 14x</td>
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<td>1968-I</td>
<td>Basold</td>
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<td>r 24</td>
<td>Silver nitrate</td>
<td>8x–19x (U) not investigated</td>
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<td>1968-II</td>
<td>Basold</td>
<td>Wheat, cv. 'Künzel'</td>
<td>r 20</td>
<td>Silver nitrate</td>
<td>6x–30x (U) not investigated</td>
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<td>1968</td>
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<td>r 2 × 6</td>
<td>Lead nitrate</td>
<td>8x–19x U 8x, 10x–12x, 16x</td>
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<td>1967</td>
<td>Basold</td>
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<td>r 4</td>
<td>Ferrosulphat</td>
<td>3x–32x U not investigated</td>
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<td>1965a</td>
<td>Boiron</td>
<td>Wheat, cv. 'Vilmorin n° 27'</td>
<td>1</td>
<td>Mercury chloride</td>
<td>3x–18x U</td>
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<td></td>
<td></td>
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<td>1963</td>
<td>Boiron</td>
<td>Wheat, cv. 'Vilmorin n° 27'</td>
<td>8x: 11; 16x: 8; 18x: 6</td>
<td>As₄Na₃H</td>
<td>3x–18x U 8x, 16x, 18x</td>
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<td>1962</td>
<td>Netien</td>
<td>Wheat, cv. 'Maitre Pierre'</td>
<td>1</td>
<td>Cobalt chloride</td>
<td>2x–18x U 8x, 12x, 15x, 18x</td>
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</tr>
</tbody>
</table>

1 Methods: b = Blinding; r = Randomisation; s = Systematic negative control experiments, as identified from the publication.
2 Controls: U = Unsuccussed potentisation medium; S = Succussed potentisation medium; P = Potentised potentisation medium; D = Diluted test substance, P-C = Positive control; N = No treatment group. Adequate controls to identify specific treatment effects (S, P) are printed in bold. Treatment effects were compared to the controls without brackets.
3 Effective potency levels: List of all potency levels which were significantly effective in any of the measured parameters, compared to the control without brackets.
4 Fertilizer content: 10% Nitrogen, 1% Ammoniacal Nitrate, 0.6% Nitrate Nitrogen, 8.4% Urea Nitrogen, 15% Phosphoric acid, 10% Potash.
on the growth of adult plants: there were two studies with duckweed,\textsuperscript{84,85} two studies with lettuce,\textsuperscript{61,79} and one study with \textit{Justicia pectoralis},\textsuperscript{52} pigeon pea\textsuperscript{86} and lady’s finger,\textsuperscript{87} respectively. The third experimental plant model used plant slips: three studies investigated the modification of the O\textsubscript{2} consumption of wheat seedling slips\textsuperscript{52,59,60} and another study investigated the growth of slips of african violet.\textsuperscript{63}

\textbf{Measured parameters:} The most commonly used experimental outcome (26 of 29 studies) was plant growth. Growth was measured by three categories: size (including shoot and root length, leaf area, leaf number or stem diameter), weight (including fresh and dry weight) and germination rate (and speed). Twelve of these 26 studies measured the effect on the plant growth by size only.\textsuperscript{53–56,62,63,75,76,84,85} Ten of the 26 studies measured further parameters: three of the ten studies\textsuperscript{52,86,87} also measured weight and secondary metabolites or biochemical parameters; two studies\textsuperscript{67,77} additionally reported weight and germination rate; three studies\textsuperscript{78} measured size and germination rate, and two studies\textsuperscript{61,79} measured size and weight. Three studies\textsuperscript{57,60} measured growth by germination rate and germination time, whereas one study\textsuperscript{53} used only weight as the measuring parameter. Besides plant growth, three of 29 studies measured the O\textsubscript{2} consumption of wheat seedlings\textsuperscript{58,59,69}.

\textbf{Experimental plant systems and treatment:} We distinguished between timing and the route of application of the treatment, to identify how the contact between plant and homeopathic remedy was effected.

\textbf{Timing:} In six studies\textsuperscript{55,56,58,59,60} the treatment of the plants was carried out before cultivation, 22 studies\textsuperscript{52–54,57,60–63,67,75–79,84,85} seedlings or plants were treated over a longer period during the cultivation, and one study\textsuperscript{73} had pre-treatment combined with treatment during cultivation.

\textbf{Route of application:} Most of the germination as well as plant slip experiments (20 studies\textsuperscript{54–60,62,63,69,73,75–78}) assured the plants contact with the homeopathic potencies by immersing them into the potencies for a certain period, mostly during the whole time of experiment. One study\textsuperscript{53} watered the plant substrate in the pots with potencies or control at the beginning of the experiments. Two other ways of treatment were the application with a spray in four studies\textsuperscript{52,79,86,87} and the daily application of one drop potency or control on the soil in one study.\textsuperscript{61} One study\textsuperscript{67} covered the seed with filter paper, soaked in potency or control, respectively. Two studies\textsuperscript{84,85} used waterplants, continuously floating in potency or control solutions.

\textbf{Potentisation:} All 29 studies with MIS $\geq 5$ were checked for potentisation technique. Eleven studies\textsuperscript{54–56,62,78,84,85} used the multiple-glass method for potentisation. Two of them\textsuperscript{55} used the single- and the multiple-glass method. In 17 studies\textsuperscript{53} no detailed information about the use of the method was given.\textsuperscript{52,53,57–61,63,67,69,73,75–77,86,87} In 15 studies\textsuperscript{53–60,62,69,84,85} the test substances were potentised by hand. About 12 different methods of hand succussion were used. Most differences were found in the number of succussion beats; beyond that, there were also differences in whether the potencies were succussed with or without hitting, or in a vertical or in horizontal movement.

Four studies\textsuperscript{67,73,75,76} used a succussion machine for potentisation. For machine succussion, essentially three different techniques were employed: in two studies,\textsuperscript{75,76} potencies were succussed for 4.5 min, in the third study\textsuperscript{73} they were succussed with 200 beats in 20 seconds, the fourth study\textsuperscript{67} potentised with 100 beats. In 10 studies\textsuperscript{52,61,63,77–79,86,87} no information was given about whether succussion was done by hand or by machine.

\textbf{Plants and potentised substances:} The germination experiment with wheat is the most commonly used experimental model: more than half of the papers with statistics (23 of 43 studies) used wheat. In the following, we have summarised all plant models that were described in the 29 studies with MIS $\geq 5$, including the studies, which investigated the specificity of homeopathic preparations (see Table 2).

\textbf{a) Wheat}

With 16 of 29 studies, wheat is the most frequently investigated plant model. Seven studies\textsuperscript{54–78} with wheat investigated the effect of Silver nitrate. With only unsuccussed potentisation medium as control, the potency levels 8x–13x (2 studies), 6x–30x, 24x and 24x–26x (2 studies) were tested. The potency levels 13x, 14x, 24x (2 studies) and 26x influenced growth of the wheat seedlings. Two of the 7 studies evaluated the effects by comparison of potency level to potency level, respectively, without factoring the control into the analysis of the results.\textsuperscript{54} Two studies\textsuperscript{54} compared silver nitrate 24x with potentised potentisation medium, and found it to be specifically effective (increase of the germination rate).

The influence of Arsenicum album potencies on the germination rate and speed of wheat seeds was investigated in three studies\textsuperscript{57,60} with a huge dataset. The first, a pilot study, tested the potency levels 23x and 30x compared to unsuccussed potentisation medium. The other two investigated the effect of 23x, 30x, 35x, 40x and 45x, and the third study additionally 27x and 42x. The second study compared the potencies to unsuccussed potentisation medium, potentised potentisation medium (H\textsubscript{2}O 30x) and diluted test substance (Arsenicum album 10–36) as control. The third study compared the potencies to unsuccussed potentisation medium. This study additionally investigated the potentised potentisation medium and the diluted test substance, both in the same potentisation levels or dilution steps as potencies used, in comparison to unsuccussed potentisation medium. The first study did not find significant potency effects. In the second study, significant differences were found compared to three sorts of control: between 25x, 30x, 40x, 45x and the unsuccussed control; between a pool of the 40x and 45x data and the potentised control (increase of the germination rate); between 25x, the 40x-45x-pool and the diluted test substance. In the third study all tested potency levels showed significant effects compared to unsuccussed control. Some of the potency levels stimulated wheat germination whereas others induced inhibition of the germination process. A multi-centre study with wheat\textsuperscript{73} investigated the specific effect of Copper sulphate 5c, 7c and 9c compared to the...
unsuccessed and the succussed potentisation medium. They tested four different ethanol-water mixture ratios of the potentisation medium and found the strongest effects (for all potency levels) with a potentisation medium with 20% ethanol.

Ferrous sulphate (3x–32x)\(^53\) and lead nitrate (8x–19x)\(^75\) were also tested with the wheat germination model. The first study evaluated potency levels only against each other. Potency levels 8x, 10x–12x and 16x of lead nitrate modified wheat growth significantly compared to unsuccessed potentisation medium. Three studies\(^{58,59,69}\) investigated the effect of homeopathic potencies on the O\(_2\) consumption of wheat seedling slips. The substances Cobalt chloride (2x–18x), Mercury chloride (3x–18x) and As\(_2\)Na\(_2\)H (3x–18x) were investigated, compared to unsuccessed potentisation medium. The O\(_2\) consumption of the seedling slips was significantly increased by the potency levels of 8x, 12x, 15x and 18x of Cobalt chloride, as well as the potency levels 9x and 10x of Mercury chloride, and the potency levels 8x, 16x an 18x of As\(_2\)Na\(_2\)H.

b) Dwarf peas

Two publications\(^{55,56}\), reporting 3 studies, investigated the specific effects of potentised plant hormones (compared to succussed and unsuccessed potentisation medium) on the growth of dwarf peas. Gibberellic acid was tested in 12x–30x, 17x, as well as 17x and 18x. Specific effects of Gibberellic acid 13x, 15x, 17x (3 studies) and 23x were found (increase of shoot length). Kinetin, Auxin and Abscisic acid were tested in the potency levels 12x–30x. Only Kinetin 19x showed a specific effect on the growth of dwarf peas (enhancing shoot length). These effects were verified by systematic negative control experiments.

c) Duckweed

Two studies\(^{84,85}\) investigated specific effects of homeopathic potencies on the growth of duckweed (Lemna gibba L.). The first study was a screening of twelve test substances in the potency levels 14x–30x, compared with successed and unsuccessed potentisation medium. In the second study experiments with four test substances used in the first study were repeated several times. Gibberellic acid, Kinetin, Silver nitrate and Lema minor were investigated in both studies. Auxin, Copper sulphate, Methyl jasmonate, Metoxuron, Phosphorus, Potassium nitrate, Sulphur and Lactose were only investigated in the first study. Silver nitrate 24x, 28x and 29x; Kinetin 14x, 16x, 20x, 23x, 26x, 27x and 30x (decrease of the growth rate) as well as Phosphorus 21x, 25x and 29x (increase of growth) showed specific effects in the screening. In the second study only Gibberellic acid 15x, 17x, 18x, 23x and 24x showed specific effects (decrease of growth). These effects were verified by systematic negative control experiments.

d) Justicia pectoralis Jacq. (L.)

One outdoor study\(^{52}\) investigated the specific effects of 8 test substances on the growth and the production of secondary metabolites (coumarine) of Justicia pectoralis Jacq. (L.). 3c potencies of Arnica montana, Justicia, Phosphorus, Sulphur, Humic acid, Acanthaceae, Cumarina P.A. and Guaco were compared with successed and unsuccessed potentisation medium; the first five preparations increased the coumarine content significantly.

e) African violet

One study\(^{63}\) investigated the specific effect of the plant hormone Indole butric acid 33x on the root and the development of new leaves of african violet slips. Enhancing root growth, the 33x showed a specific effect, compared to the successed potentisation medium.

f) Lettuce (Lactura sativa)

One outdoor study\(^{66}\) investigated the specific effects of Silicea 1LM and Sulphur 1LM on the growth of lettuce. For both test substances specific effects were found (decrease of weight and breadth), in comparison to the successed potentisation medium. Another outdoor study\(^{79}\) investigated the effect of Carbo vegetabilis (6c, 12c, 30c, 100c, 200c) on the growth of lettuce, compared with unsuccessed potentisation medium and an untreated control group. All potency levels were found effective.

g) Mung seed

One publication\(^{61}\) investigated the effects of the potency levels 1x–24x of a fertilizer on the growth of mung seedlings. Regrettably, the author only reported that the potencies were effective (compared to unsuccessed potentisation medium, the diluted test substance and a positive control), but no detailed information about the potency levels was given.

h) Sida rhombifolia

One study\(^{64}\) investigated the effect of Cymbopogon winterianus Jowitt (Citronella) in the potency levels 3c, 6c, 12c, 24c and 30c on the germination and growth of Sida rhombifolia. The potency levels 6c, 12c and 30c showed significant effects (in comparison to the unsuccessed control) on the parameter root and shoot length, fresh weight, germination rate, germination time and speed. The 3c was found effective on the root and shoot length, and the 24c influenced the root length and the fresh weight significantly.

i) Pigeon pea and

j) Lady’s finger

Two outdoor studies\(^{86,87}\) investigated the specific effects of potencies of two growth retardants normally used in agriculture on multiple parameters of growth and plant biochemistry. The potency levels 30c and 200c of CCC (2-chloroethyl) trimethyl ammonium chloride), CCC (nano) and MH (Malec hydradize) were tested in pigeon pea and lady’s finger. Growth and biochemical parameters of both plants were significantly influenced (increase) by the potencies of CCC and CCC (nano) and MH.

Tested substances: In the 29 studies with MIS ≥ 5 the effect of 32 different test substances was investigated. Twenty-two studies investigated only one test substance. Five studies examined two to four test substances. Moreover, there were two screenings, one study with eight different test substances and another study with twelve.

The most frequently used test substances were Silver nitrate (nine studies), Gibberellic acid (five studies), and Kinetin and Sulphur in three studies each (See also Table 2, column Tested substances).
Table 3a  This table includes all 15 studies (with MIS ≥ 5) using adequate controls to investigate specific effects of homeopathic preparations. Sub-table 3a lists the 6 studies consisting of only one independent experiment, sub-table 3b all 9 studies with multiple experiments. The table lists plant species, experimental methods, number of plants per treatment and experiment, number of independent experiments, measured parameters, way of treatment, potentised substances, potentisation method, tested potency levels, controls, statistical tests and effective potency levels.

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<th>Author &amp; year</th>
<th>Reference</th>
<th>Plant</th>
<th>Methods</th>
<th>Number n (per treatment and experiment)</th>
<th>Number of independent experiments</th>
<th>Measured parameters</th>
<th>Treatment</th>
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<tr>
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<td>87</td>
<td>Lady's finger (Abelmoschus esculentus L. Moench)</td>
<td>10 plants</td>
<td>1</td>
<td>Length, girth and weight of shoots; length and weight of roots; number of leaves/plant; weight, area and water content of leaves; cholinphyl and protein content</td>
<td>D: 1:500 in water diluted potencies, with a spray, 2 times daily, only the first 2 days of exp.</td>
<td>CCC, CCC (nano), MH</td>
<td>H: Mother tinctures: Tribinations of CCC, MH, CCC+Copper nano particles with lactose; Potentiation: 1:100 dilution steps in Alcohol 90%, Succussion: 10 beats</td>
<td>30c, 200c</td>
<td>P</td>
<td>One way ANOVA, Hest</td>
<td>30c, 200c (all test substances)</td>
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<tr>
<td>Sukul 2008</td>
<td>86</td>
<td>Pigeon pea (Cajanus cajan L. Millsp.)</td>
<td>20 plants</td>
<td>1</td>
<td>Leaves- and branches number, shoot girth at day 75; shoot weight, root length, number of flowers/plant at day 150; cholinphyl-, carbohydrate- and protein-content of leaves</td>
<td>D: with a spray on several days of exp., 1:500 in water diluted potencies</td>
<td>CCC (2-chloroethyltrimethylammonium chloride), CCC (nano), MH (Maleic hydrazide)</td>
<td>H: Mother tinctures: 1g CCC/MH per 1 ml Alcohol 90%; 1 g CCC+Copper nano particles triturated with lactose per 1 ml Alcohol 90%; Potentiation: 1:100 dilution steps in Alcohol 90%, Succussion: 10 beats</td>
<td>30c, 200c</td>
<td>P</td>
<td>One way ANOVA, Hest</td>
<td>30c, 200c (all test substances)</td>
<td></td>
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<td>Scherr 2007</td>
<td>84</td>
<td>Duckweed (Lemna gibba L.)</td>
<td>b; r; s 5 beakers per parameter</td>
<td>1</td>
<td>Frond area and frond number related growth rate, day 0–7, 0–3, 3–7</td>
<td>D: The waterplants grew in potency or control</td>
<td>Silver nitrate, Copper sulphate, Gibberelllic acid, Auchen, Kinetin, Ladoine, Lemna minor, Methyl Jasmonate, Metoxuron, Potassium nitrate, Sulphur, Silica, Globuli (Haitos Homeopathic Pharmacy) solved in dist. water</td>
<td>H: horizontal, no hitting; multiple-glass method</td>
<td>14x—30x</td>
<td>U + S</td>
<td>With F-test protected Fisher’s LSD</td>
<td>Arg nitr: 24x, 28x, 29x; Kinetin: 14x, 16x, 20x, 23x, 26x, 27x, 30x; Phos: 21x, 25x, 29x</td>
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<td>Chapman 2004</td>
<td>61</td>
<td>Lettuce (Lactuca sativa), cv. ‘Tom Thumb’</td>
<td>b; r</td>
<td>17–25 plants</td>
<td>1</td>
<td>Plant height, weight, bread, 55 days after germination</td>
<td>D: 1 drop on the soil, daily</td>
<td>Silver nitrate, Copper sulphate, Gibberelllic acid, Kinetin, Ladoine, Lemna minor, Methyl Jasmonate, Metoxuron, Potassium nitrate, Sulphur, Silica</td>
<td>Globuli (Haitos Homeopathic Pharmacy) solved in dist. water</td>
<td>LM1</td>
<td>S</td>
<td>Student's Hest</td>
<td>LM1</td>
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<td>Andrade 2001</td>
<td>52</td>
<td>Justicia pectoralis L. (Jacqu.)</td>
<td>b; r</td>
<td>16 plants</td>
<td>1</td>
<td>Fresh and dry weight, leaf area, coumarine content</td>
<td>D: weekly application with a spray</td>
<td>Justice, Acanthaceae, Cumarine P.A.; Guaco; Phosphos; Arnicamontana; Humic acid</td>
<td>Hahnemannian, no detailed information</td>
<td>3c</td>
<td>(U); P</td>
<td>Scott-Knott test, Hest</td>
<td>Justicia; Phos; Sulphur; Arnicamontana; Humic acid: 3c</td>
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<td>Pongratz 1994a-II</td>
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<td>Wheat, cv. ‘Mephisto’</td>
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<td>200 seeds</td>
<td>1</td>
<td>Stalk length, germination rate, after 5 days</td>
<td>D: Immersion of the seed</td>
<td>Silver nitrate</td>
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<td>24x</td>
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<td>Chi-square test; one way ANOVA</td>
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<tr>
<td>Scherr 2009</td>
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<td>Duckweed (Lemna gibba L.)</td>
<td>b; r; s 5 beakers</td>
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<td></td>
<td>Gibberelllic acid, Kinetin, Silver nitrate, Lemna minor</td>
<td>H: horizontal, no hitting; multiple-glass method</td>
<td>14x—30x</td>
<td>U + S</td>
<td>With F-test protected Fisher’s LSD</td>
<td>Gibberelllic acid: 15x, 17x, 18x, 23x, 24x</td>
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<tr>
<td>Reference</td>
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<tr>
<td>Baumgartner 2008</td>
<td>2008</td>
<td>Dwarf peas (Pisum sativum L.), cv. 'Frueher Zwerg'</td>
<td>b; r; s 50 seedlings</td>
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<td>Shoot length after 14 days. B: Immersion of the seed. H: vertical, no hitting; multiple-glass method</td>
<td>17x, 18x U + S</td>
<td>With F-test protected Fisher's LSD</td>
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<td>Baumgartner 2004-i</td>
<td>2004</td>
<td>Dwarf peas (Pisum sativum L.), cv. 'Frueher Zwerg'</td>
<td>b; r; s 23 seedlings</td>
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<td>Brizzi 2000</td>
<td>2000</td>
<td>Wheat (Triticum durum L.), cv. 'MEC'</td>
<td>b; r 33 seeds per treatment</td>
<td>unclear</td>
<td>Germination rate, germination speed. D: Watering at the beginning of the experiment. H: vigorous hitting, 70 impacts</td>
<td>23x, 25x, 27x, 30x, 35x, 40x, 45x U; (P; D)</td>
<td>Poisson test, global poison comparison test, odds ratio</td>
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<td>Betti 1994-II</td>
<td>1994</td>
<td>Wheat (Triticum durum L.), cv. 'MEC'</td>
<td>b; r 99 seeds</td>
<td>8</td>
<td>Germination rate, germination speed. D: Watering at the beginning of the experiment. H: vigorous hitting, 70 impacts</td>
<td>23x, 25x, 30x, 35x, 40x, 45x U; (D)</td>
<td>(U; D) Poisson test, parametric one-factor ANOVA, Chisquare test,</td>
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<td>Endler 1991</td>
<td>1991</td>
<td>African violet (Saintpaulia sp.)</td>
<td>b; r 7574 slips per experiment</td>
<td>3</td>
<td>Rooting, development of new leaves after several weeks. D: Immersion of the plant slips. H: serial dilution, succussion after each dilution step.</td>
<td>33x S</td>
<td>Not parametric U-tests (Mann-Whitney)</td>
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<td>Pongratz 1990</td>
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<td>Wheat, cv. 'Mephisto'</td>
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<td>10</td>
<td>Stem length, dry weight of shoots and roots, after 5 days; germination rate. D: Immersion of the seed. H: serial dilution, succussion after each dilution step.</td>
<td>24x P</td>
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<td>Noiret 1979b</td>
<td>1979</td>
<td>Wheat, cv. 'Hardi'</td>
<td>b unclear</td>
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<td>Fresh and dried weight after 4-5 days of germination. B and D: Immersion of the seed. H: serial dilution, succussion after each dilution step.</td>
<td>5c, 7c, 9c U; S</td>
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</tbody>
</table>

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1 Methods: b = Blinding; r = Randomisation; s = Systematic negative control experiments, as identified from the publication.
2 Treatment: B = Before cultivation; D = During cultivation.
3 Potentisation: H = Hand succussion; M = Machine succussion.
4 Controls: U = Unsuccessed potentisation medium; S = Successed potentisation medium (adequate control); P = Potentised potentisation medium (adequate control); D = Diluted test substance, P-C = Positive control. Treatment effects were compared to the controls without brackets.
5 Effective potency levels: List of all potency levels which were significantly effective in any of the measured parameters, compared to the control without brackets.
Studies with controls adequate to investigate specific effects of homeopathic preparations: Fifteen studies included adequate controls to identify specific effects of the tested homeopathic potencies (Table 3). Twenty-five test substances were investigated with eight plants: wheat, dwarf peas, duckweed, lettuce, African violet, Justicia pectoralis, pigeon pea and lady’s finger (See Table 3, columns Plants, Tested substances and Controls).

The most frequently tested potentised substances were Gibberellic acid (5 studies), Silver nitrate (4 studies), Kinetin and Sulphur (3 studies each), as well as Arsenicum album. Auxin, Phosphorus, Copper sulphate, Lemna minor, CCC, CCC(nano) and Maleic hydrazide in 2 studies each. The following test substances were investigated in one study only: Abscisic acid, Acanthaceae, Arnica montana, Coumarine P.A., Guaco, Indole butric acid, Humic acid, Justicia, Lactose, Methyl jasmonate, Metoxuron, Potassium nitrate and Silicea (see Table 4).

Four studies55,63,77,78 investigated the effect of only one potency level (17x; 24x (2 studies); 33x). All four studies found the tested potencies to be significantly effective. Another study61 investigated one potency level (1LM) of two test substances, and both significantly influenced the growth of the plants. One study73 tested three potency levels (5c, 7c and 9c) of one substance: all potency levels induced significant effects. Furthermore, two studies86,87 investigated two potency levels (30c, 200c) of three test substances; significant effects of all potency levels and test substances were observed.

In a screening, one study52 tested the potency level 3c of eight different test substances. Five of the potentised substances showed specific effects. The phenomenon that

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<th>Effect†</th>
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<td>Dwarf peas</td>
<td>12x–30x; 17x</td>
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<td>I</td>
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<td>Dwarf peas</td>
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<td>Dwarf peas</td>
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<td>17x</td>
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<tr>
<td></td>
<td></td>
<td>Duckweed</td>
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<td>Silver nitrate</td>
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<td>Wheat</td>
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<td>Wheat</td>
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<td>77</td>
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<tr>
<td></td>
<td></td>
<td>Duckweed</td>
<td>14x–30x</td>
<td>24x; 28x; 29x</td>
<td>D</td>
<td>84</td>
</tr>
<tr>
<td>Kinetin</td>
<td>3</td>
<td>Dwarf peas</td>
<td>12x–30x</td>
<td>19x</td>
<td>I</td>
<td>55</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Duckweed</td>
<td>14x–30x</td>
<td>14x, 16x, 20x, 23x, 26x, 27x, 30x</td>
<td>D</td>
<td>84</td>
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<tr>
<td></td>
<td></td>
<td>Duckweed</td>
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<td></td>
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</tr>
<tr>
<td></td>
<td></td>
<td>Lettuce</td>
<td>1LM</td>
<td></td>
<td></td>
<td>61</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Duckweed</td>
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<td>No effect</td>
<td></td>
<td>84</td>
</tr>
<tr>
<td>Arsenicum album</td>
<td>2</td>
<td>Wheat</td>
<td>23x, 30x; 35x, 40x, 45x</td>
<td>40x + 45x</td>
<td>I</td>
<td>57</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wheat</td>
<td>23x, 27x, 30x; 35x, 40x, 42, 45x</td>
<td>42x, 45x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Auxin (IAA)</td>
<td>2</td>
<td>Dwarf peas</td>
<td>12x–30x</td>
<td>No effect</td>
<td></td>
<td>55</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Duckweed</td>
<td>14x–30x</td>
<td>No effect</td>
<td></td>
<td>84</td>
</tr>
<tr>
<td>Copper sulfate</td>
<td>2</td>
<td>Duckweed</td>
<td>14x–30x</td>
<td>No effect</td>
<td></td>
<td>84</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Duckweed</td>
<td>14x–30x</td>
<td>No effect</td>
<td></td>
<td>85</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Duckweed</td>
<td>14x–30x</td>
<td>No effect</td>
<td></td>
<td>84</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wheat</td>
<td>5c, 7c, 9c</td>
<td>5c, 7c, 9c</td>
<td>D</td>
<td>73</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lady’s finger</td>
<td>30c, 200c</td>
<td></td>
<td></td>
<td>87</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lady’s finger</td>
<td>30c, 200c</td>
<td></td>
<td></td>
<td>87</td>
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<tr>
<td>Maleic hydrazide</td>
<td>2</td>
<td>Pigeon pea</td>
<td>30c, 200c</td>
<td>30c, 200c</td>
<td>I</td>
<td>86</td>
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<tr>
<td></td>
<td></td>
<td>Pigeon pea</td>
<td>30c, 200c</td>
<td>30c, 200c</td>
<td>I</td>
<td>86</td>
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<td>30c, 200c</td>
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<td>Lady’s finger</td>
<td>30c, 200c</td>
<td></td>
<td></td>
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<tr>
<td>Phosphorus</td>
<td>2</td>
<td>Justicia pectoralis</td>
<td>3c</td>
<td>3c</td>
<td>I</td>
<td>52</td>
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<tr>
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<td>Duckweed:</td>
<td>14x–30x</td>
<td>21x, 25x, 29x</td>
<td>D</td>
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</tr>
<tr>
<td>Ascisic acid</td>
<td>1</td>
<td>Dwarf peas</td>
<td>12x–30x</td>
<td>No effect</td>
<td></td>
<td>55</td>
</tr>
<tr>
<td>Acanthaceae</td>
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<td>Justicia pectoralis</td>
<td>3c</td>
<td>No effect</td>
<td></td>
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</tr>
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<td>Arnica montana</td>
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<td>I</td>
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<tr>
<td>Coumarine P.A.</td>
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<td>No effect</td>
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<tr>
<td>Guaco</td>
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<td>Justicia pectoralis</td>
<td>3c</td>
<td>No effect</td>
<td></td>
<td>52</td>
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<td>Indole butric acid</td>
<td>1</td>
<td>African violet</td>
<td>33x</td>
<td>33x</td>
<td>I</td>
<td>63</td>
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<td>Humic acid</td>
<td>1</td>
<td>Justicia pectoralis</td>
<td>3c</td>
<td>3c</td>
<td>I</td>
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<tr>
<td>Justicia</td>
<td>1</td>
<td>Justicia pectoralis</td>
<td>3c</td>
<td>3c</td>
<td>I</td>
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<td>Lactose</td>
<td>1</td>
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<td>14x–30x</td>
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<tr>
<td>Methyl jasmonate</td>
<td>1</td>
<td>Duckweed</td>
<td>14x–30x</td>
<td>No effect</td>
<td></td>
<td>84</td>
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<tr>
<td>Meloxuron</td>
<td>1</td>
<td>Duckweed</td>
<td>14x–30x</td>
<td>No effect</td>
<td></td>
<td>84</td>
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<tr>
<td>Potassium nitrate</td>
<td>1</td>
<td>Duckweed</td>
<td>14x–30x</td>
<td>No effect</td>
<td></td>
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<tr>
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<td>Lettuce</td>
<td>1LM</td>
<td></td>
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</table>

† I = Increase; D = Decrease.

Table 4 This table gives an overview of the 15 studies investigating specific effects, focussing on the tested substances. It shows how many investigations were done with which potentised substance, which plant models were used, which potency levels were investigated and which potency levels showed specific effects.
only some test substances influenced plant growth was also found in the second screening, where specific effects were observed only for three of twelve potentised substances.54,84

Some of the experiments,55,56,77,78 investigating only one or few potency levels were based on preceding studies with potency level series.77,78,85 In these experiments, only some potency levels seemed to be effective. That only some or few potency levels seem to be biologically active was observed in nearly all studies investigating specific effects of series of potencies.55,57,60,84,85 Three of these studies verified their results by systematic negative control experiments; therefore this phenomenon does not seem to be an artefact.

When similar potency levels of the same substance were tested with two different plant organisms, some potency levels showed effects for both plants, e.g. Gibberellic acid in dwarf peas and in duckweed.85 The tested intersecting potency levels were 14x–30x, and concordant effective potency levels were 15x, 17x and 23x, which increased growth for dwarf peas and decreased for duckweed. Another example are potencies of Silver nitrate with wheat (two studies77,78) and with duckweed.84 In all three studies the 24x was found to be effective.

Two studies – one with dwarf peas,56 the other with duckweed85 – found significant interactions between the treatment and the experiment number. This means that the effects were not identical for the different experiments. The results of both studies were verified by systematic negative control experiments, so this effect does not seem to be due to instabilities of the experimental systems. A further phenomenon is the opposite effect of several potency levels of the same substance in one study.60 Two potency levels (27x and 35x) of Arsenicum album inhibited wheat germination, whilst 30x, 40x, 42x and 45x stimulated it.

In five studies which used unsuccussed and succussed potentisation medium as control,55,56,84,85 no significant difference between succussed and unsuccussed control was found. In two further studies,57,60 however, when unsuccussed and potentised potentisation medium were used as control, H2O 30x and 45x (i.e. succussed control) induced a significant modification of wheat germination.

Reproducibility: There are only very few investigations of the external reproducibility in the group of studies with MIS  = 5 and most of them repeated experiments of Kolisko27 (published 1926), whose results were evaluated without statistical analysis. Two studies54,76 which investigated the effect of Argentum nitricum (8x–19x) on the growth of wheat seedlings both found significant effects, but the effective potency levels were not identical. One study78 investigated the effects of three potency levels of a series (Argentum nitricum 24x–26x), and found the same phenomenon as Kolisko, 24x and 26x were more effective than 25x. Additionally, a multi-centre study73 with four participating laboratories investigated the effect of Copper sulphate 5c, 7c and 9c, on the germination of wheat seeds. Effects were mostly similar, but there was imprecision in potency production between the laboratories.

Internal reproducibility was investigated by five55,56,84,85 studies, three with dwarf peas and two with duckweed, which included all of the quality criteria (see Table 2, column 3). In these studies, a significant interaction between treatment and experiment number was observed for some test substances; the effects of certain potency levels varied between inhibiting, inducing and no modification of plant growth. To determine conditions for successful reproducibility, one study56 with dwarf peas investigated Gibberellic acid 17x and 18x, using four different harvest lots of the same seed cultivar. Significant effects on plant growth were found only for Gibberellic acid 17x for two of the four seed harvests, and Gibberellic acid 18x did not induce any significant growth modification. Different results in carbohydrate content for the four pea seed batches indicated that differing seed quality could cause problems in the reproducibility of the results in experimental plant systems.

A further study86,87 repeated the investigation of 3 potentised plant retardants, maintaining the set-up, but replacing the plant model. Here, the test substances were effective in many parameters of the two plant models, but the effects were not always the same.

Discussion

Almost all the studies observed effects of homeopathic potencies on plants, even in high dilutions far beyond the Avogadro number. It is conspicuous that no linear or monotonous relationship between potency level and effect size was observed in any of the studies that tested a series of potency levels. Consistently across all studies, only some of the potency levels of a series were found to be effective, and not all potency levels were effective in the same way. In three studies55,84,85 investigating potency series from 12x or 14x to 30x, a minimum of one potency level and a maximum of seven potency levels of the series showed significant effects. Another study found some of the tested potency levels stimulated germination, but other potency levels inhibited it.60 There was no uniform activity of all potency levels of a continuous series. This observation is not an artefact due to multiple statistical testing, since three studies55,84,85 used the protected Fisher’s LSD-Test with predefined statistical hypotheses. Another study60 used a corresponding statistical procedure with an exact global Poisson test preceding comparisons of single potency levels to the control. Both procedures minimize type 1 as well as type 2 errors. In addition, stability of the experimental system (including the statistical procedure applied) was ensured by systematic negative control experiments in three studies.

The observation that only some potency levels of a tested series of potencies were effective was not only observed in plant studies. A multi-centre study on the effect of potentised histamine on human basophil cells88 also showed differing effective potency levels of potency series. In addition, Linde89 found in his “Critical review and meta-analysis of serial agitated dilutions in experimental toxicology” that Mercury 15c induced a 40% mortality reduction in mercury-intoxicated mice, whilst Mercury 9c was ineffective. Thus the phenomenon of alternating levels of active and inactive potencies seems to be of a general intrinsic
nature among homeopathic potencies in preclinical investigations. Possible consequences of this observation for clinical application should be studied more closely. For example, in case of lack of success of a well-indicated remedy, it might be worth trying other potencies.

One aim of this review was to identify studies that provide evidence for specific effects of homeopathic remedies, i.e. effects implying some sort of “memory” of the carrier substance (e.g. water) for the mother tincture diluted. We identified 15 studies which included adequate (succussed) controls. In these cases, the results obtained most probably cannot be attributed to non-specific effects, such as molecules and ions dissolved from the potentisation vessel during the succussion process. Five studies compared the succussed and unsuccussed potentisation medium statistically and found no significant difference between these controls. This may mean that plants in general are not influenced by non-specific succussion effects. If this were true, studies with unsuccussed controls may also be indicative of specific remedy effects. We think, however, that empirical confirmation is needed to prove that this is truly the case for any plant system investigated.

Several studies reported effects of potentised plant growth substances or plant hormones. Since even stronger responses were observed in basic research animal models after application of potentised animal hormones, one might setup the hypothesis, that human hormones – unknown at Hahnemann’s times – might be promising substances for human homeopathic therapy, corresponding homeopathic drug provings might be very interesting.

In two studies, series of potencies and analogous unsuccussed dilutions of the test substances were investigated. One study found significant effects of the potencies in comparison to the diluted controls, and the other study found significant effects for most of the tested potency levels, but no effects of any dilution level of the diluted test substance. If these systems are not sensitive to non-specific succussion effects, these results imply that succussion is a necessary part of the homeopathic remedy production procedure. It thus would be interesting to conduct more investigations involving different sorts of controls to determine the relevance of the non-specific processes during succussion for plant models, to assess the usefulness of the different controls, and determine what intensity or sort of succussion is necessary to yield effective homeopathic preparations.

Some of the studies with specific effects had shortcomings in the quality of the experimental set-up; for example, blinding and randomisation were missing. Moreover, we could not assess the standardisation of the laboratory or ambient conditions during the experiments in most of the publications. Only five studies documented the stability of the experimental system by publishing data about systematic negative control experiments. In our opinion, this is the only way of convincingly demonstrating the stability of the experimental set-up chosen, i.e. that there were no false-positive effects caused by influences of laboratory or ambient conditions. This is a very important point for the quality of a study, but systematic negative control experiments have not so far been implemented on a routine basis. Especially in research in homeopathy, where in many cases effects are not explicable on molecular level, distinguishing effects from artefacts is essential. One example of the situation, where the instability of the experimental system precluded interpretation of the results, was an extensive investigation about the effect of potentised ambient pollutants (metals) on wheat germination. In this case, the systematic negative control experiments yielded evidence for instability of the experimental set-up. We assume that many of the researchers carried out systematic negative control experiments to prove the stability of their experimental plant system. Without a proper description of the results of these control experiments, however, it is difficult to assess the validity of the study results definitively.

We found only a few published studies that tried to reproduce earlier findings, and these studies were only partially successful. There may be an unknown number of unpublished experiments reporting unsuccessful attempts to reproduce other trials. The lack of reproduction trials points to problems with reproducibility. One study started to investigate causes for reproduction problems in homeopathic basic research, and hypothesized differing seed quality as a limiting factor. These results have some similarity to the presumption, that basophil cells of different human donors vary in their sensitivity to treatment with potentised histamine.

The studies reviewed were conducted for different reasons. Some studies can be interpreted as homeopathic drug provings on plants, corresponding to human homeopathic drug provings as introduced by Hahnemann. The aim of other studies was to look for alternative fertilisation methods for plant production. Further studies strived to develop simple test systems for providing scientific evidence of the specificity of homeopathic remedies, to generate tools to control homeopathic drug production, and to ensure product quality.

Using plant models for research in homeopathy offers several advantages such as studies with large data sets, with a short experimental running time, and avoidance of the placebo effect as well as the ethical problems of trials on animals or humans. The multiplicity of the reviewed studies and the diversity of the results reflect the complexity of the issue, however. In homeopathic drug provings with humans, a large variety of symptoms are depicted to understand the homeopathic effects of a potentised substance. Most homeopathic studies with healthy plants use only one or few measuring parameters, such as growth, length, weight, area or germination rate. This limitation could be one of the reasons why only few of many test substances were found to be effective in screening studies. To deepen the understanding of homeopathic effects on plants, more sophisticated research would be interesting. For example, plant growth could be measured by multiple methods, or parameters mirroring the plant’s physiology and biochemistry could be used, as they are in some newer studies.

Many different potentisation techniques were used in the studies reviewed: there were considerable variations in the intensity and movement of succussion, for example. Most studies identified effects of the tested potencies, but to
date we have not learned how the different potentisation techniques influence the efficacy of the potentised agents. To establish a closer relationship between homeopathic basic research and human and veterinary homeopathic care, it would make sense to investigate potencies, which were produced by the established procedures also used by homeopathic pharmacies.

Some studies investigated similar series of potency levels of the same test substance – using different plant models in some cases – but they started potentisation with different concentrations of the mother tinctures. As a result, when we examine the possibility that there could be particular potency levels of one test substance which are effective in multiple plant models, we cannot say whether these potency levels (e.g. the 24x of Argentum nitricum, used in two studies) are comparable because of their differing mother tincture concentrations.

One aim of this review was to identify useful models for future studies in homeopathic basic research. The study model should be simple and easily transferable to other laboratories. Moreover, the standard deviation should be as small as possible. The germination model with wheat has been the most frequently used model so far. In spite of a rather high standard deviation it seems to be sensitive enough to identify potency effects. Treating the seeds with liquid immersion is simple. Parameters to measure growth modifications are root and shoot length, fresh and dry weight, and germination rate and speed. Wheat is routinely used for studies with poisoned or stressed plants as well as for studies with healthy plants. Wheat germination studies with information about systematic negative control experiments are still lacking, however.

The study model based on duckweed has generated attention. Duckweeds are small monocotyledonous waterplants with predominantly vegetative growth that are used as standard test organisms in ecotoxicology. Due to the uniform plant growth the duckweed studies benefited from a very low standard deviation, though the effects were also quite small. The waterplants grew directly in potency or control, with parameters being the area and number of the fronds, measured using a scanner with an image processing system.

Using systematic negative control experiments, the reliability of the system was assessed over the entire course of investigations. The homogeneity of plant growth, the option of simple treatment and the standardised bioassay all qualify the duckweed model as a very interesting test system. The installation of the measuring technique involves considerable effort and expenditures. Easier and cost-efficient techniques may be developed in future to make this bioassay easily suited for multi-centre use.

Other interesting experimental plant models use the concentration of plant secondary metabolites or other biochemical substances as outcome parameters. This allows depicting a physiological reaction of the plants to the treatment with homeopathic potencies. These studies demand established techniques for analysis, special laboratory equipment and corresponding experience; this may be challenging for future independent repetitions in other laboratories.

Conclusions

Healthy plant models seem to be a useful tool to investigate basic research questions about the specificity of homeopathic preparations. Homeopathic basic research could move forward by conducting plant studies of high quality design that includes systematic negative control experiments, blinding, randomisation, adequate statistical analysis, and appropriate controls to identify specific remedy effects. It would be advisable to do more trials on the potentisation process itself, and to use standardised potentisation techniques so that studies can be easily compared. Only few studies attempted independent reproduction trials. Increased cooperation between active laboratories would be advisable for identifying the crucial parameters for successful reproduction trials.

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