Systematic review and statistical analysis of the integrity of 33 randomized controlled trials

ABSTRACT

Background: Statistical techniques can investigate data integrity in randomized controlled trials (RCTs). We systematically reviewed and analyzed all human RCTs undertaken by a group of researchers, about which concerns have been raised.

Methods: We compared observed distributions of p values for between-groups differences in baseline variables, for standardized sample means for continuous baseline variables, and for differences in treatment group participant numbers with the expected distributions. We assessed productivity, recruitment rates, outcome data, textual consistency, and ethical oversight.

Results: The researchers were remarkably productive, publishing 33 RCTs over 15 years involving large numbers of older patients with substantial comorbidity, recruited over very short periods. Treatment groups were improbably similar. The distribution of p values for differences in baseline characteristics differed markedly from the expected uniform distribution ($p = 5.2 \times 10^{-32}$). The distribution of standardized sample means for baseline continuous variables and the differences between participant numbers in randomized groups also differed markedly from the expected distributions ($p = 4.3 \times 10^{-4}, p = 1.5 \times 10^{-5}$, respectively). Outcomes were remarkably positive, with very low mortality and study withdrawals despite substantial comorbidity. There were very large reductions in hip fracture incidence, regardless of intervention (relative risk 0.22, 95% confidence interval 0.15–0.31, $p < 0.0001$, range of relative risk 0.10–0.33), that greatly exceed those reported in meta-analyses of other trials. There were multiple examples of inconsistencies between and within trials, errors in reported data, misleading text, duplicated data and text, and uncertainties about ethical oversight.

Conclusions: A systematic approach using statistical techniques to assess randomization outcomes can evaluate data integrity, in this case suggesting these RCT results may be unreliable.

GLOSSARY

RCT = randomized controlled trial.

Investigating concerns about the integrity of data from biomedical research is difficult. One approach is to use statistical techniques to compare the observed distributions of baseline variables from a group of randomized controlled trials (RCTs) for which concerns about data integrity exist, with the expected distributions that would arise if treatment allocation occurred by chance. This approach was previously used to identify a case of fraudulent data involving at least 168 RCTs.

Meta-analysts may identify concerns about data integrity during their careful review of large numbers of publications. While undertaking systematic reviews in osteoporosis, we considered RCTs by Yoshihiro Sato et al. ("the researchers") for inclusion. This group of authors has published a very large number of RCTs (appendix e-1, table e-1A, references A1–A33 at Neurology.org) that collectively have substantially influenced relevant systematic reviews. Dr. Sato is the first author on 79% of these trials and holds primary oversight of the majority of these publications. Questions had been raised in journal correspondence regarding some of these papers, and we had additional concerns about aspects of a number of papers identified from the Department of Medicine (M.J.B., G.D.G., A.G.), University of Auckland, New Zealand; and Health Services Research Unit (A.A.), University of Aberdeen, Foresterhill, Scotland.

Go to Neurology.org for full disclosures. Funding information and disclosures deemed relevant by the authors, if any, are provided at the end of the article.
for our reviews. We wondered whether the statistical techniques previously used for comparing distributions of baseline variables could be applied to a systematic review of RCTs from the researchers that included 3 papers published in Neurology® (A9, A17, A23, which have been retracted; for a full list of papers in this study retracted to date, see table e-1B.). In addition, we assessed some other potentially relevant aspects of the body of RCTs including the researchers’ productivity and recruitment rates, outcome data, textual consistency within and between RCTs, and ethical oversight.

METHODS In November 2012, we searched MEDLINE and Embase for all RCTs with Y. Sato or Yoshihiro Sato listed as an author, published in English or Japanese. Two hundred fifty-three potentially relevant publications were identified. After excluding unrelated articles and articles by other individuals with the same name, we identified 33 RCTs in humans (table e-1A). For each report, we extracted data on study design, ethical approval, randomization, baseline characteristics, and outcomes, and identified inconsistencies in reporting of methods and results.

Halberek et al.4 suggested that “baseline data indicate that the study groups are strikingly well matched” in many RCTs reported by the researchers,5 who responded that this occurred “not by design but by chance.” Since allocation of participants in an RCT is random, comparisons between randomized groups for independent variables at baseline should produce a uniform distribution of p values; for example, there is an equal likelihood of a p value of <0.1 and >0.9, of <0.2 and >0.8, etc. Baseline variables within an RCT are not always independent and may be correlated. However, this interdependence is unlikely to translate into substantial between-group similarities where simple randomization is used. Clustering of the p values for differences between groups may not occur in the same direction or to the same degree in other trials. Since any differences between groups are attributable to chance, if there is consistent clustering of variables in multiple trials, the distribution of p values across multiple trials from a group of clustered variables should still be approximately uniform. Therefore, across a body of RCTs, clustering is unlikely to affect the distribution of p values markedly. We compared the observed distribution of p values in deciles for all baseline comparisons with the expected distribution using the χ² test (Excel 2007; Microsoft, Redmond, WA). We used reported p values from the papers, and when these were not reported, we calculated p values from the reported summary data (mean, SD, or percentage) using t tests or 1-way analysis of variance for continuous variables and χ² test or Fisher exact test for categorical variables (OpenEpi version 2.3.1, www.OpenEpi.com).

We undertook a complementary analysis to assess the sampling distribution of continuous variables. The central limit theorem states that if a population is repeatedly sampled, the means of these samples are approximately normally distributed. The mean of the sample means (x̄) approximates the population mean (μ) and the SD of the sample means is the standard error of the mean (which is the population SD/√n). Any normal distribution curve can be standardized to produce a curve with mean 0 and SD 1 using the formula (x − μ)/SD. A randomized group in an RCT can be considered as a random sample from the entire trial population. We calculated the trial population mean and SD for each baseline continuous variable from the summary data presented in the trial report. We then selected the control group (or the first presented treatment group when only active treatments were compared) and standardized the sample mean (x − μ)/SEM. We tested whether the distribution of these standardized sample means differed from the expected distribution (standardized SD of 1) using a t test for the equality of variance (Excel 2007), in an approach similar to that used by Carlisle.6

We assembled a dataset of 13 RCTs6-20 conducted by the Auckland Bone and Joint Research Group (table e-2) as a matched control group of trials that were similarly sized to the RCTs of interest, conducted over a similar time frame, and also focus on the prevention or treatment of osteoporosis in older people. We compared the distributions of p values and standardized sample means between this control dataset and the dataset of interest. Bootstrap resampling was used to account for ignorance of the dependence structure between differences in baseline variables and single group variance structures. We randomly selected between 350 and 450 baseline p values (with replacement) and between 250 and 350 baseline standardized sample means (with replacement) from both datasets, performed 2-sample Kolmogorov–Smirnov tests on these values (SAS v9.4; SAS Institute Inc., Cary, NC), and repeated this 1,000 times.

In a trial with simple randomization, the proportion of participants in each group is determined by chance, and thus the numbers of participants in the randomized groups in a set of trials will be binomially distributed.21,22 For a 2-arm trial with 2 participants, the probability of equal number of participants in each group (i.e., one) is 50%, for 4 participants is 37.5%, for 10 participants is 25%, and for 50 participants is 11%. For block randomization, the number of participants in each trial arm is equal in each block. Thus, any differences between the size of randomized groups arise from the last block. If the block is filled, the groups will be equal in size. For a block size of 2 or 4 in a 2-arm trial, an odd number of total participants means the groups differ by one. For a block size of 4, 2 participants in the final block gives a 2/3 probability the groups are equal in size, and a 1/3 probability that they differ by 2 participants.22 When block randomization is stratified, the same principle applies for each stratum. We used the approach of Carlisle2 to assess whether the distribution of differences in numbers of participants between treatment groups in the trials conducted by the researchers is consistent with the binomial distribution.

Hip fracture was the most frequently reported fracture outcome, reported for 23 trials. Using random-effects models, we pooled these outcome data from trials by the researchers and compared the results to published contemporaneous meta-analyses of trials from other investigators. Statistical heterogeneity between trial-level summary data was assessed using the Cochran Q statistic (p < 0.10) and the P statistic (P > 50). Analyses were performed using Comprehensive Meta-Analysis version 2 (Biostat, Englewood, NJ).

RESULTS Productivity and recruitment rates. Table 1 shows the characteristics of the 33 human RCTs (A1–A33). Of note, these 33 RCTs were published over a 15-year period and included 5,894 participants. Three (A9, A17, A23) were published in Neurology. Y. Sato was the first author for 26 RCTs. Another author was the first author for the remaining 7 RCTs and a coauthor for 25 RCTs, and 3 others coauthored at least 10 RCTs. The rate of
publication of RCTs suggests a very large well-funded research network, with access to a very large catchment of patients with substantial comorbidity who, nonetheless, are willing trial participants. Considering these practical requirements, it was surprising that there was no statement regarding funding for 30 of these practical requirements, it was surprising that there was no statement regarding funding for 30 of the 33 human RCTs. Furthermore, none of the RCTs contained a statement on trial registration, although many trials were published before this became customary.

The human RCTs represent remarkable productivity, particularly given the frailty of the participants in several of the trials. For example, in the 5 months between March 2003 and July 2003, the researchers recruited 500 ambulatory female patients older than 70 years with Alzheimer disease living in the

### Table 1  Study design of 33 human randomized controlled trials by the researchers

<table>
<thead>
<tr>
<th>Ref</th>
<th>Year</th>
<th>Patient group</th>
<th>Age</th>
<th>Sex</th>
<th>Intervention</th>
<th>Endpoint</th>
<th>Duration</th>
<th>No. Follow-up</th>
<th>Recruitment period</th>
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<tr>
<td>A1</td>
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<td>Stroke</td>
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<td>ACD vs P</td>
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<td>84</td>
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<td>Stroke</td>
<td>NIl</td>
<td></td>
<td>Either</td>
<td>Vit K vs nil</td>
<td>BMD</td>
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<td>108</td>
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<td>NIl</td>
<td></td>
<td>Either</td>
<td>Ipri vs ACD vs nil</td>
<td>BMD</td>
<td>12 mo</td>
<td>103</td>
</tr>
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<td>ACD vs P</td>
<td>BMD</td>
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<td>86</td>
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<td>BMD</td>
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<td>120</td>
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<td>Stroke</td>
<td></td>
<td></td>
<td>Either</td>
<td>B12 vs folate vs both homocysteine</td>
<td>BMD</td>
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<td>BMD</td>
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<td>NIl</td>
<td></td>
<td>Female</td>
<td>Etid vs P</td>
<td>Bone metabolism</td>
<td>3 mo</td>
<td>80</td>
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<td>NIl</td>
<td></td>
<td>Female</td>
<td>Etid vs Alend</td>
<td>BMD</td>
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<td></td>
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<td>Median NCS</td>
<td>2 y</td>
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<td></td>
<td>Either</td>
<td>B12/folate vs P</td>
<td>Hip fracture</td>
<td>2 y</td>
<td>628</td>
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<td>Stroke</td>
<td>NIl</td>
<td></td>
<td>Female</td>
<td>Vit D vs P</td>
<td>Falls</td>
<td>2 y</td>
<td>96</td>
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<td>&gt;65</td>
<td></td>
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<td>Hip fracture</td>
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<td>BMD</td>
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<td>Stroke</td>
<td>&gt;65</td>
<td></td>
<td>Female</td>
<td>Rised vs P</td>
<td>Hip fracture</td>
<td>12 mo</td>
<td>374</td>
</tr>
<tr>
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<td>AD</td>
<td>&gt;70</td>
<td></td>
<td>Female</td>
<td>Rised vs P</td>
<td>Fracture</td>
<td>18 mo</td>
<td>500</td>
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<td>AD</td>
<td>&gt;70</td>
<td></td>
<td>Female</td>
<td>Vit K/vit D/Ca vs nil</td>
<td>Osteoporosis</td>
<td>2 y</td>
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<td>&gt;55</td>
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<td>BMD</td>
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<td>50</td>
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<td>NIl</td>
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<td>Etid vs P</td>
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<td>82</td>
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<td></td>
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<td>Alend vs P</td>
<td>Hip fracture</td>
<td>2 y</td>
<td>288</td>
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<tr>
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<td>PD</td>
<td>&gt;65</td>
<td></td>
<td>Male</td>
<td>Rised vs P</td>
<td>Osteoporosis</td>
<td>2 y</td>
<td>242</td>
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<tr>
<td>A24</td>
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<td>NIl</td>
<td></td>
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<td>Alend vs Ral</td>
<td>BMD</td>
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<tr>
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<td>2009</td>
<td>Volunteers</td>
<td>&gt;50</td>
<td></td>
<td>Either</td>
<td>Exercise vs nil</td>
<td>Falls</td>
<td>5 mo</td>
<td>68</td>
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<tr>
<td>A28</td>
<td>2011</td>
<td>Osteoporosis</td>
<td>NIl</td>
<td></td>
<td>Female</td>
<td>Alend vs Elca</td>
<td>Back pain</td>
<td>6 mo</td>
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<td>A29</td>
<td>2010</td>
<td>AD</td>
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<td></td>
<td>Either</td>
<td>Vit K vs nil</td>
<td>Hip fracture</td>
<td>12 mo</td>
<td>231</td>
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<td>A30</td>
<td>2011</td>
<td>PD</td>
<td>NIl</td>
<td></td>
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<td>Hip fracture</td>
<td>2 y</td>
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<td>PD</td>
<td>&gt;65</td>
<td></td>
<td>Either</td>
<td>Sunlight vs nil</td>
<td>Hip fracture</td>
<td>2 y</td>
<td>324</td>
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<td>A32</td>
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<td>Stroke</td>
<td>NIl</td>
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<td>Female</td>
<td>Vibration vs nil</td>
<td>Physical function</td>
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</table>

**Abbreviations:** ACD = alphacalcidol; AD = Alzheimer disease; Age = age restriction; Alend = alendronate; ALS = amyotrophic lateral sclerosis; B12 = vitamin B12; BMD = bone mineral density; Ca = calcium; Elca = elcatonin; Etid = etidronate; Ipri = ipriflavone; Methylpred = methylprednisolone; NCS = nerve conduction studies; NMS = neuroleptic malignant syndrome; NS = not stated; P = placebo; PD = Parkinson disease; Ral = raloxifene; Ref = reference; Rised = risedronate; Vit = vitamin.

*5 trials actively recruiting participants, and b3 trials with active follow-up between March and July 2003.
community in 2 months (A18), 280 male patients older than 65 years with hemiplegic stroke in 2 months (A15), and 374 female patients older than 65 years with acute hemiplegic stroke in 4 months (A17). Compounding the workload of recruiting for these 3 trials, participants were also reviewed every 4 weeks, and during this period, there was also ongoing intensive follow-up in 3 trials involving 774 participants (A13, A14, A20) and recruitment and intensive follow-up for 2 other trials involving 292 participants (A11, A23). Despite the high workload required to conduct clinical trials on such a scale, 4 of these 8 trials listed only the same 4 coauthors (A14, A15, A17, A18), and only one additional coauthor was listed for 2 of these trials (A13, A23). However, the recent Journal of Bone and Mineral Research retraction notice for one paper (A16) states that the coauthors of Dr. Sato “are named as such for honorary reasons and are not responsible for the content of the manuscript.” Two of the 3 papers in Neurology reported details on potential participant eligibility. In one, <10% of potential participants were ineligible despite very restrictive inclusion criteria (A9). In both, >85% of people approached were enrolled (A9, A23).

Neuroleptic malignant syndrome can rarely occur following reduction or withdrawal of levodopa therapy. The researchers identified 40 patients with Parkinson disease with this syndrome over 3 years in a single institution, Futase Social Insurance Hospital, which has only 55 inpatient beds and 4 clinicians (http://futase-hp.jp/about/outline.php) (A8). A correspondent found this astonishing because clinicians at their institution, which has a special interest in Parkinson disease, could only “recall 2 such cases in living memory.” Correspondents also expressed disbelief at the recruitment rate in the female stroke trial (A17) because, in their experience, <10% of stroke admissions would meet the eligibility criteria.

**Improbably similar randomized treatment groups.** In 32 human RCTs, baseline data for all participants in all randomized groups were presented for 513 variables; data were only presented for study completers in one RCT (A1). Figure 1A shows the distribution of \( p \) values for between-groups comparisons of these 513 variables. Of note, 52% of \( p \) values were >0.8, while only 6%, 14%, and 27% of \( p \) values were <0.2, <0.4, and <0.6, respectively. This distribution is highly unlikely to have arisen by chance (\( p = 5.2 \times 10^{-43} \)). In the 25 RCTs with Y. Sato as the first author, the distribution of \( p \) values differed significantly from the expected distribution (\( p = 3.8 \times 10^{-100} \)) (figure 1C). Figure 1B shows the distribution of the standardized sample means for 402 baseline continuous variables. The SD of these standardized sample means was 0.84, and the distribution differed markedly from the expected distribution (\( p = 4.3 \times 10^{-5} \)), with values clustered more tightly around the mean than expected. Figure 1D shows that in the 25 RCTs with Y. Sato as the first author, the distribution of standardized sample means differed from the expected distribution with values clustered tightly around the mean (\( p = 1.5 \times 10^{-13} \)). The distribution of \( p \) values from 918 baseline variables from 13 RCTs in the control dataset was consistent with the expected independent uniform distribution (\( p = 0.07 \), figure e-1), and the SD of the standardized sample means from 726 baseline variables was 1.02, also consistent with the expected distribution (\( p = 0.78 \)). However, comparison of baseline \( p \) values and standardized sample means from the RCTs of interest and the control RCTs using bootstrap resampling showed marked differences in distributions (baseline \( p \) values: \( p < 0.001 \) in all 1,000 comparisons; standardized sample means: median \( p \) value 6 \( \times 10^{-4} \), 95% confidence interval 2 \( \times 10^{-7} \) to 0.021).

Table 2 shows that the numbers of participants in each treatment group were the same in 27 of 30 two-arm RCTs. For all 33 trials, randomization in blocks was stated specifically for 10 RCTs, using computer-generated random numbers for 13 RCTs, no details were reported for 9 RCTs, and one trial was pseudorandomized using alternate allocation. Four trials that used block randomization were stratified by site, and for one trial, participant numbers by site were reported in correspondence about the paper (A18). For another 2 trials (A13, A15), stratified randomization was not described, but multiple sites and participant numbers by site were reported in corrections or correspondence about the papers. We assumed randomization was stratified by site for these 2 trials. Figure 2 shows that the distribution of differences in participant numbers between the treatment groups differs from the expected distribution (\( p = 5 \times 10^{-39} \)) in 20 two-arm RCTs presumed to use simple randomization (i.e., unless block randomization was stated). For the 10 RCTs that used block randomization, the observed distribution of differences was consistent with the expected distribution (\( p = 0.48 \)). When all 30 two-arm trials were considered, the observed distribution of differences differed from the expected distribution (\( p = 1.5 \times 10^{-10} \)). Finally, we treated all trials as if they used randomization with a block size of 4 unless another approach was explicitly stated (except trial A1, which has a difference of 6 participants between groups meaning that block randomization is very unlikely). The distribution of differences in participant numbers between the treatment groups still differs from the expected
Figure 1  Distribution of p values and standardized sample means in 32 randomized controlled trials by the researchers

A. All variables: n=513

\[ p = 5.2 \times 10^{-32} \]

B. All variables: n=402, mean -0.02, SD 0.84

\[ p = 4.3 \times 10^{-4} \]

C. Trials by Sato: n=399

\[ p = 3.8 \times 10^{-100} \]

D. Trials by Sato: n=311, mean -0.059, SD 0.68

\[ p = 1.5 \times 10^{-11} \]

E. Study A13: n=23

F. Study A13: n=13, mean -0.23, SD 0.37

G. Study A23: n=12

H. Study A23: n=10, mean -0.10, SD 0.54

(A, C, E, G) The observed vs expected distribution of p values by decile in 32 randomized controlled trials for all baseline variables (A), in 25 trials with Y. Sato as first author (C), and as illustrative examples, for all baseline variables in study A13,
analyses of other trials for these agents,24 substantially greater than those observed in meta-
also shows that the magnitude of these reductions is
the relatively large numbers of agents and the diver-
This lack of heterogeneity is very surprising given
the relatively large numbers of agents and the diver-
Despite studying frail elderly individuals with sub-
stantial comorbidity, there were consistent and sub-
stantial reductions in hip fracture incidence regardless of the treatment studied (relative
risk 0.22, 95% confidence interval 0.15–0.31, \( p < 0.0001 \), range of relative risk 0.10–0.33) with very
little statistical heterogeneity between the results.
This lack of heterogeneity is very surprising given
the relatively large numbers of agents and the diver-
Figure 3 shows the observed reductions in hip
fracture in RCTs with a control group that received
either placebo or no treatment for the various treat-
ments studied by the researchers. There were consis-
tent and substantial reductions in hip fracture
incidence regardless of the treatment studied (relative
risk 0.22, 95% confidence interval 0.15–0.31, \( p < 0.0001 \), range of relative risk 0.10–0.33) with very
little statistical heterogeneity between the results.
This lack of heterogeneity is very surprising given
the relatively large numbers of agents and the diver-
ity of the populations studied. Furthermore, figure 3
also shows that the magnitude of these reductions is
substantially greater than those observed in meta-
analyses of other trials for these agents,24–26 and the confidence intervals around the pooled result for the
researchers’ trials do not overlap the confidence intervals
around the pooled results for trials conducted by
other investigators. Thus, the very positive results of
the trials by the researchers are inconsistent with re-
sults from other groups.

Logical and other inconsistencies. There are a number of inconsistent results between trials conducted by
the researchers. For example, in a trial in Alzheimer
disease (A29), the control group received risedronate
and calcium and had a hip fracture rate of 86/1,000
patient-years. Yet, the active treatment arm of another
trial in patients with Alzheimer disease (A18) who
received risedronate, calcium, and vitamin D had
a hip fracture rate of 15/1,000 patient-years. Similarly,
trials in Parkinson disease in which the control group received placebo plus vitamin D
reported very high rates of hip fracture (49, 37, and
55/1,000 patient-years) (A22, A23, A30) but when
sunlight exposure was used as the active treatment
arm to improve vitamin D status, the hip fracture
rate was only 9/1,000 patient-years (A31). Likewise,
when vitamin D was used in the placebo arm of trials,
bone density decreased substantially by 0.9% to 3.2%
(A18, A22, A23, A30), whereas when sunlight
exposure was studied as active treatment, bone
mineral density increased by 2.7% to 3.8% (A16, A31).

There are also inconsistencies within trials. For
example, in one Neurology paper (A9), the authors
state that 109 participants were randomized to one
group and 108 to the other. However, the particip-
ant flow diagram shows 129 participants in each
group with 109 and 108, respectively, completing the
trial. In one study (A22) it is stated that random-
ization was performed by one of the authors “(J.I.),”
and in the same paragraph, it is stated that “follow-
up assessment of patient’s condition was performed
by physicians (Y.S., J.I.) who did not participate in
the initial randomization.” In one study (A29), it is
stated that randomization was stratified by site, yet
only one site was mentioned in the methods section.
In another study (A15), participants were eligible for
inclusion if they had sustained a stroke at least 3
months before the study began, but the mean dura-
tion of illness at baseline in both randomized groups
was 90 days, or slightly less than 3 months, which
appears implausible.

There are also errors in the reported outcome
data. In 14 RCTs, rates of hip fracture per 1,000
patient-years are reported (table 2). For 11 of these
trials, the reported rates can simply be calculated
with the equation: rate = number with hip frac-
ture/(participants \( \times \) duration of trial). These
rates are not correct because they do not account for loss
to follow-up or censoring of participants who have
sustained a hip fracture. Data on adverse events are
also misreported. For example, in 2 of the 3

Figure 1 legend, continued:
the subject of the recent expression of concern from the Journal of the American Medical Association27 (E), and study A23, published in Neurology (G). The dotted line shows the expected proportion (0.1) for each decile. (B, D, F, H) The observed vs
expected distribution of standardized sample means for all continuous baseline variables in these 32 trials (B) and in 25
trials with Y. Sato as first author (D), and for all continuous baseline variables in study A13 (F) and study A23 (H). The
expected distribution is a normal distribution curve with mean 0 and SD 1 (dotted line). The solid line is a normal curve of
best fit to the observed data. All the graphs show that the observed distribution differs markedly from the expected
distribution.
<table>
<thead>
<tr>
<th>Study</th>
<th>Participants</th>
<th>Method of randomization</th>
<th>Hip fracture</th>
<th>Withdrawals</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No.</td>
<td></td>
<td>No.</td>
<td>Rate (per 1,000 patient-y)</td>
</tr>
<tr>
<td></td>
<td>Control/Treat</td>
<td></td>
<td>Control/Treat</td>
<td>Control/Treat</td>
</tr>
<tr>
<td>A1</td>
<td>39/45</td>
<td>Not described</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>A2</td>
<td>54/54</td>
<td>Not described</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>A3</td>
<td>34/34/35</td>
<td>Not described</td>
<td>1</td>
<td>0/0</td>
</tr>
<tr>
<td>A4</td>
<td>43/43</td>
<td>CG random no.</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>A5</td>
<td>49/49</td>
<td>CG random no.</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>A6</td>
<td>60/60</td>
<td>CG random no.</td>
<td>8</td>
<td>1</td>
</tr>
<tr>
<td>A7</td>
<td>63/64/64*</td>
<td>CG random no.</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>A8</td>
<td>20/20</td>
<td>CG random no.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A9</td>
<td>129/129</td>
<td>CG random no.</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>A10</td>
<td>40/40</td>
<td>CG random no.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A11</td>
<td>25/25*</td>
<td>Alternating</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>A12</td>
<td>68/67</td>
<td>CG random no.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A13</td>
<td>314/314</td>
<td>CG random no., permuted blocks of 4</td>
<td>27</td>
<td>6</td>
</tr>
<tr>
<td>A14</td>
<td>48/48</td>
<td>CG random no., permuted blocks of 4</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>A15</td>
<td>140/140</td>
<td>CG random no., permuted blocks of 4</td>
<td>10</td>
<td>2</td>
</tr>
<tr>
<td>A16</td>
<td>132/132</td>
<td>CG random no.</td>
<td>9</td>
<td>2</td>
</tr>
<tr>
<td>A17</td>
<td>197/197</td>
<td>CG random no.</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>A18</td>
<td>250/250</td>
<td>CG random no., strat. permuted blocks of 4</td>
<td>19</td>
<td>5</td>
</tr>
<tr>
<td>A19</td>
<td>100/100</td>
<td>CG random no.</td>
<td>15</td>
<td>2</td>
</tr>
<tr>
<td>A20</td>
<td>25/25</td>
<td>Not described</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A21</td>
<td>41/41</td>
<td>CG random no., strat. permuted blocks of 4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A22</td>
<td>144/144</td>
<td>CG random no., permuted blocks of 4</td>
<td>14</td>
<td>4</td>
</tr>
<tr>
<td>A23</td>
<td>121/121</td>
<td>CG random no., permuted blocks of 4</td>
<td>9</td>
<td>3</td>
</tr>
<tr>
<td>A24</td>
<td>61/61*</td>
<td>Not described</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>A25</td>
<td>56/56*</td>
<td>Not described</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A26</td>
<td>34/34</td>
<td>Not described</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A27</td>
<td>40/40</td>
<td>CG random no., strat. permuted blocks of 2</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>A28</td>
<td>97/97*</td>
<td>Not described</td>
<td>NA</td>
<td>0/0</td>
</tr>
<tr>
<td>A29</td>
<td>115/116</td>
<td>CG random no., strat. permuted blocks of 4</td>
<td>10</td>
<td>2</td>
</tr>
<tr>
<td>A30</td>
<td>136/136</td>
<td>CG random no., permuted blocks of 4</td>
<td>15</td>
<td>3</td>
</tr>
<tr>
<td>A31</td>
<td>162/162</td>
<td>CG random no.</td>
<td>11</td>
<td>3</td>
</tr>
<tr>
<td>A32</td>
<td>41/41*</td>
<td>CG random no.</td>
<td>NA</td>
<td>1/0</td>
</tr>
<tr>
<td>A33</td>
<td>26/26</td>
<td>Not described</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: CG = computer generated; NA = not applicable as there was no control group; strat. = stratified by site. Blanks indicate data not reported by the researchers. *Indicates active treatment comparison studies.
Neurology papers, it was stated that no adverse events (A17) or no serious adverse events (A23) occurred during the trial. However in A17, 10 participants withdrew because of death or intercurrent illness, 87 experienced at least one fall, and 8 had a hip fracture. Likewise, in A23, 7 participants withdrew because of death or intercurrent illness, 12 had a hip fracture, and there was an average of 1.5 falls per participant during the trial.

Misleading text. In one study (A13), the researchers claimed to have recruited 628 trial participants from a single site. In a later correction, they stated that there were 3 additional unnamed hospitals. The researchers stated that the reason for this misinformation was that “these hospitals were reluctant to have their names in the article.” Given the large size and intensive nature of these trials, and the absence of statements regarding funding, we presume the trials were effectively funded by the hospitals involved, with substantial trial input from other unnamed hospital staff. It is hard to understand why the hospital and its staff would not accept any recognition of their substantial contributions.

Duplicate data and text. There are examples of strikingly similar text in the reports, including duplicate outcome data. Table 3 shows that the reported mean and SD for fall rates were identical in 3 studies (A4, A6, A9) for both the treatment and control groups, and another study (A19) had identical SD to the earlier studies but the mean differed by 1 unit for both the treatment and control groups. Later, these data were altered without explanation in corrections for 2 of these trials (A6, A9). Duplicated text is also apparent when logical errors arise. Table 3 shows an example of 2 studies with identical exclusion criteria, but one criterion does not make sense for one trial because it only applies to women, but only men were eligible to participate in the trial. Some examples of duplicated text are extensive. The majority of the text in one article (A16) is identical or nearly identical to that of a later article (A31) (table e-3).

Ethical oversight. In the 33 RCTs, the ethics committee that approved the research was usually listed as the institutional or local ethics committee (22 studies). By using all the published information regarding the ethics committee approval and the location of the study, it appears that ethical approval was granted by the ethics committee of Futase Social Insurance Hospital in 11 RCTs, Mitate Hospital for 10 RCTs, Keiyu Orthopaedic Hospital for 5 RCTs, multiple committees in 2 RCTs, and in 5 RCTs the committee could not be identified. We searched the internet for “ethics” and “Mitate Hospital” or “Futase” and were unable to identify any studies performed by other groups approved by either of these ethics committees. With only 4 clinicians practicing at Futase Hospital, it seems likely that members of the Futase Hospital ethics committee had at least some clinical relationships with at least some of the coauthors of the papers by the researchers. In fact, in at least 6 studies (A9, A16, A17, A21, A27, A31), the chairperson of the ethics committee appeared to be involved with the with stroke and 500 patients with Alzheimer disease, respectively, from a single site. In a later correction, they stated that there were 2 additional unnamed hospitals. The researchers stated that the reason for this misinformation was that “these hospitals were reluctant to have their names in the article.” Given the large size and intensive nature of these trials, and the absence of statements regarding funding, we presume the trials were effectively funded by the hospitals involved, with substantial trial input from other unnamed hospital staff. It is hard to understand why the hospital and its staff would not accept any recognition of their substantial contributions.

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conduct of the study as this person is reported to have performed the randomization and allocation of participants to treatment groups.

DISCUSSION Our systematic review and application of established methods to examine data integrity to the 33 human RCTs published by the researchers raises serious concerns about the integrity and validity of the reported results. Objective evidence for these concerns is provided by the statistical analyses, which demonstrate a systematic failure of randomization, and the consistently outlying outcome data when...
### Table 3: Examples of duplication of text in articles by the researchers

<table>
<thead>
<tr>
<th>Article</th>
<th>Text</th>
<th>Article</th>
<th>Comparative text</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>A4</td>
<td>Falls rate (mean, SD); treatment group 1.3 (1.9), control group 1.4 (1.8)</td>
<td>A6, A9</td>
<td>Treatment group 1.3 (1.9), control group 1.4 (1.8)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>A19</td>
<td>Treatment group 2.3 (1.9), control group 2.4 (1.8)</td>
<td></td>
</tr>
<tr>
<td>A22</td>
<td>Patients with impairment of renal, hepatic, cardiac, or thyroid function or those who had known causes of osteoporosis, such as primary hyperparathyroidism or renal osteodystrophy, were excluded from this study. Patients were excluded if they had been treated with corticosteroids, estrogen, calcitonin, bisphosphonate, calcium, or vitamins D and K for 3 months or more during the 12 mo preceding the study, and those who had been administered these agents for even a brief period during the preceding 2 mo were also excluded.</td>
<td>A23</td>
<td>Patients with impairment of renal, hepatic, cardiac, or thyroid function or those who had known causes of osteoporosis, such as primary hyperparathyroidism or renal osteodystrophy, were excluded from this study; Patients were excluded if they had been treated with corticosteroids, estrogen, calcitonin, bisphosphonate, calcium, or vitamins D and K for 3 mo or more during the 12 mo preceding the study, and those who had been administered these agents for even a brief period during the preceding 2 mo were also excluded.</td>
<td>Trial A23 was restricted to men. The exclusion criterion regarding estrogen treatment does not make sense in context of elderly males with Parkinson disease.</td>
</tr>
<tr>
<td>A16</td>
<td></td>
<td>A31</td>
<td>The highlighted text in A16 is identical or nearly identical to the text in A31 (table e-3, included with permission from Wiley).</td>
<td></td>
</tr>
</tbody>
</table>

Compared to those from other researchers. Taken together with the implausible productivity of the group, internal inconsistencies for outcome data in their work, duplication of data, numerous misleading statements and errors, and concerns regarding ethical oversight, our analysis suggests that the results of at least some of these trials are not reliable. Recently, the *Journal of the American Medical Association* has issued an expression of concern about one trial (A13), and the *Journal of Bone and Mineral Research* an expression of concern followed by retraction of another trial (A16), both precipitated by our concerns.²³⁻²⁸

Two issues arise from our analyses of distribution of $p$ values and standardized sample means for considering the success or failure of randomization. There will be differences in $p$ values and standardized sample means calculated from published summary data compared to values calculated from raw data because of rounding of variables. However, any differences would be small and could not explain the marked departure from the expected distributions that we found. A second issue is that some of the baseline variables may not be independent. For example, if by chance, one randomized group is older than the other, this group may have similar differences in variables correlated with age, such as bone mineral density. This could lead to clustering of $p$ values and standardized means within an individual RCT. However, clustering may not occur in the same direction or magnitude in other trials. Since any differences between groups are attributed to chance, consistent clustering of variables should still produce an approximately uniform distribution of $p$ values of between-groups comparisons from a group of clustered baseline variables across multiple trials. Therefore, clustering of results from individual RCTs is unlikely to affect the analysis of a body of RCTs markedly. However, results from individual RCTs or small differences from expected variations for a body of RCTs should be interpreted cautiously.

Previously, Carlisle analyzed the distribution of standardized sample means from 168 RCTs published by an individual author, the integrity of whose work had been questioned, along with data from 366 RCTs from other authors.² He reported that the observed distribution of standardized sample means for weight, height, and age from the 366 control RCTs were consistent with expected (range of standardized SD 0.93–1.06), whereas the distributions of sample means from the 168 RCTs of interest were not (range SD 0.55–0.62). Similarly, pooling all continuous variables produced a distribution similar to expected for the 366 RCTs from other authors, whereas the distribution was markedly different to expected for the variables from the 168 RCTs of interest.² The results of our analyses of the data from the researchers were similar to the results of Carlisle’s analysis, with observed distributions of sample means markedly different to expected. A limitation of this work, and the previous work by Carlisle, is that the statistical theory underpinning the methods used has not been fully developed.

In summary, statistical techniques, such as those used in this report and similar approaches used previously by Carlisle,² that involve comparisons of the observed distributions of baseline $p$ values or baseline variables in RCTs with the expected distributions may be useful when considering or investigating the integrity of a group of RCTs.

**AUTHOR CONTRIBUTIONS**

Mark Bolland: drafting/revising manuscript, study concept or design, analysis or interpretation of data, acquisition of data, statistical analysis.

Alison Avenell: drafting/revising manuscript, study concept or design, analysis or interpretation of data. Greg Gamble: drafting/revising manuscript, study concept or design, analysis or interpretation of data.
statistical analysis. Andrew Grey: drafting/revising manuscript, study concept or design, analysis or interpretation of data. Dr. Bolland has full responsibility for the data, the analyses and interpretation, and the conduct of the research, has full access to all of the data, and has the right to publish any and all data, separate and apart from the guidance of any sponsor.

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DISCLOSURE

The authors report no disclosures relevant to the manuscript. Go to Neurology.org for full disclosures.

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REFERENCES


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