Handbook of Biomedical Research Writing: The Results and Discussion Section

by Jocelyn Graf
Hanyang University
Center for Teaching and Learning
English Writing Lab
jocelyngraf@gmail.com
http://ctl.hanyang.ac.kr/writing

Version 1
March 2008

This is a self-study guide written to help graduate students and researchers at Hanyang University in Seoul, Korea, write for publication in English in biomedical and life sciences. Parts may be helpful to other non-native speaking scientists writing in English as well as other novice researchers. Any part may be freely distributed in electronic or print form for non-commercial, educational purposes either for self-study or classroom use as long as information indicating attribution is included as follows:

Some rights reserved 2008 Jocelyn Graf and HYU CTL http://ctl.hanyang.ac.kr/writing/

However, if you want to pass the text on to others, please have them download this file from the original URL rather than passing it on as an attachment. See

http://hanyangowl.org

For more information on appropriate use see

http://creativecommons.org/licenses/by-nc-sa/2.0/kr/deed.en_US

For a definition of non-commercial use see
http://ocw.mit.edu/OcwWeb/web/terms/terms/index.htm#noncomm

This edition is currently being revised. Please email suggestions or corrections to jocelyngraf@gmail.com and adamturner7@gmail.com
4.0 The Results and Discussion Sections

The results and discussion sections present your research findings and your analysis of those findings. A few papers also contain a conclusion section, which usually focuses on practical application or provides a short summary of the paper. The results, discussion and conclusion sections are combined into one chapter in this book because they are sometimes combined in journal articles. Most articles do not contain all three sections.

4.1 The Purpose of the Results and Discussion Sections

To review, the traditional journal article in the sciences consists of four parts: Introduction, Methods, Results, and Discussion/Conclusion (IMRD). They answer these questions:

- Why do we care about the problem and the results?
- What problem are you trying to solve?
- How did you go about solving or making progress on the problem?
- What's the answer?
- What are the implications of your answer?

The last two questions are the object of the results and discussion sections, respectively. If a paper contains a conclusions section, it also focuses on implications.

4.2 The Structure of the Results Section

The *Annals of Internal Medicine*’s Information for Authors provides the following advice for preparing the results section of a clinical journal article:

> Fully describe the study sample so that readers can gauge how well the study findings apply to their patients (external validity). Then present primary findings followed by any secondary and subgroup findings. Use tables and figures to demonstrate main characteristics of participants and major findings. Avoid redundancy between text and tables and figures.


In clinical medicine, there are a number of organizations of scholars that have developed regulations for reporting on various types of studies. Before drafting your article, even if you plan to publish in a smaller journal, check the author’s guidelines of a major journal for a list of these recommendations. The *Annals of Internal Medicine* and *BMJ* have excellent detailed author’s guidelines and links to checklists. For example, the following is an excerpt from the checklist for studies of diagnostic accuracy. The group of scholars is named “STARD.”

**STARD checklist for reporting of studies of diagnostic accuracy: Results section**

<table>
<thead>
<tr>
<th>Section and Topic</th>
<th>Information to Be Included</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RESULTS</strong></td>
<td></td>
</tr>
<tr>
<td>Participants</td>
<td>When study was performed, including beginning and end dates of recruitment.</td>
</tr>
<tr>
<td></td>
<td>Clinical and demographic characteristics of the study population (at least information on age, gender, spectrum of presenting symptoms).</td>
</tr>
<tr>
<td></td>
<td>The number of participants satisfying the criteria for inclusion who did or did not undergo the index tests and/or the reference standard; describe why participants failed to undergo either test (a flow diagram is strongly recommended).</td>
</tr>
<tr>
<td>Test results</td>
<td>Time-interval between the index tests and the reference standard, and any treatment administered in between.</td>
</tr>
</tbody>
</table>
Distribution of severity of disease (define criteria) in those with the target condition; other diagnoses in participants without the target condition.

A cross tabulation of the results of the index tests (including indeterminate and missing results) by the results of the reference standard; for continuous results, the distribution of the test results by the results of the reference standard.

Any adverse events from performing the index tests or the reference standard.

Estimates

- Estimates of diagnostic accuracy and measures of statistical uncertainty (e.g. 95% confidence intervals).
- How indeterminate results, missing data and outliers of the index tests were handled.
- Estimates of variability of diagnostic accuracy between subgroups of participants, readers or centers, if done.
- Estimates of test reproducibility, if done.


Biomedical researchers doing basic science can follow a more flexible structure, depending on the nature of the study and the journal. The sample description for experimental papers would typically appear in the methods section, not the results, as mentioned above for clinical papers. Start with the most important finding, and continue through each result in a logical way. This may be according to time, if one result followed from the next, or it may be from most to least important. An example is provided later in this chapter, in the section entitled “Results Structure of an Experimental Study in the IRD(m) Format.”

4.3 The Structure of the Discussion Section

According to the Instructions for Authors of the Journal of the American Medical Association (JAMA), the discussion section is “a comment section placing the results in context with the published literature and addressing study limitations.”


Similarly, Cell’s Instructions for Authors stipulates that “the Discussion should explain the significance of the results and place them into a broader context. It should not be redundant with the Results section. This section may contain subheadings and can in some cases be combined with the Results section.”


Interestingly, because Cell does not have a clinical focus, the role of discussing limitations is not central enough to mention in the Instructions for Authors. In fact, unlike clinical journals, journals like Cell that publish long experimental reports are more likely to allow combining the results and discussion into a single section and may not contain a conclusion section at all.

Turner notes that the discussion section is a mirror image of the introduction. While the introduction starts with general background information and moves to the specific purpose of the author’s research, the discussion starts with an analysis of the author’s own specific results and moves to general implications of the research.


Tip
The Annals of Internal Medicine journal’s Information for Authors offers these recommendations for structuring the discussion section:

1. Provide a brief synopsis of key findings, with particular emphasis on how the findings add to the body of pertinent knowledge.
2. Discuss possible mechanisms and explanations for the findings.
3. Compare study results with relevant findings from other published work. Briefly state literature search sources and methods (e.g., English-language MEDLINE search to Jan 2007) that identified previous pertinent work. Use tables and figures to help summarize previous work when possible.
4. Discuss the limitations of the present study and any methods used to minimize or compensate for those limitations.
5. Mention any crucial future research directions.
6. Conclude with a brief section that summarizes in a straightforward and circumspect manner the clinical implications of the work.


4.4 Examples of Results and Discussion Sections

4.4.1 Clinical Study Results and Discussion Structure with Traditional IMRD Format

Below is presented most of the text from the results and discussion sections of a clinical report on the relationship between eating meat and getting various types of cancer.


Note: It was possible to reprint long excerpts of this journal article thanks to the PLoS Medicine journal's generous open-access policy. For more details, see http://journals.plos.org/plosmedicine/license.php.

<table>
<thead>
<tr>
<th>Comments</th>
<th>Results Section</th>
</tr>
</thead>
<tbody>
<tr>
<td>The first sentence of the results section summarizes the procedure detailed previously in the methods section. A summary of the findings for the first variable—red meat intake—is provided.</td>
<td>During a mean follow-up of 6.8 y, 53,396 cancer diagnoses (36,907 male cases and 16,489 female cases) were ascertained. The mean energy–adjusted red meat intake in this cohort was 34.6 g/1,000 kcal (38.0 g/1,000 kcal in men and 29.5 g/1,000 kcal in women). The medians of extreme quintiles ranged from 9.8 to 62.7 g/1,000 kcal for red meat and 1.6 to 22.6 g/1,000 kcal for processed meat.</td>
</tr>
<tr>
<td>The characteristics of patients studied are related to the amount of meat that they eat.</td>
<td>In general, those in the highest quintile of red meat intake tended to be . . . [demographic characteristics, habits and health] (Table 1).</td>
</tr>
</tbody>
</table>
| The first subheading. Within this section, results for the first of two types of meat are presented. First: A list of all the types of cancer risk increased by eating red meat. Next: statistical procedures that clarified the connection between meat and cancer was not related to other causes. | Red Meat

Individuals in the highest quintile of red meat intake, compared with those in lowest, had a statistically significant elevated risk of several malignancies (Table 2), including esophageal, . . . colorectal, . . . liver, . . . lung, and borderline statistical significance for laryngeal cancer . . . The positive association for red meat intake and colorectal cancer was due more to cancer of the rectum . . . than the colon, . . . Additional fine control for smoking did not alter the associations for cancers of the esophagus, colorectum, liver, lung, or larynx. In addition, the tests for interaction between smoking and both red meat . . . and processed meat . . .
intake for lung cancer risk were not statistically significant. . .

<table>
<thead>
<tr>
<th>A list of cancer types not related to eating red meat.</th>
<th>Red meat intake was not associated with gastric or bladder cancer, leukemia, lymphoma, or melanoma. The associations between red meat and cancer are summarized in Figure 1 . . .; the figure also shows the null findings for sex-specific cancers, such as . . .</th>
</tr>
</thead>
</table>
| Reference to a figure that contains the data of the first two paragraphs. | Unexpectedly, red meat intake was inversely associated with endometrial cancer. . .

**Unexpected findings. Red meat seems to help prevent one kind of cancer.**

<table>
<thead>
<tr>
<th>In further sex-specific analyses, red meat intake was positively associated with pancreatic cancer among men only . . . We observed no differences in risk by sex . . .</th>
</tr>
</thead>
<tbody>
<tr>
<td>Findings that were relevant only for men in the study. Mention of no difference between women and men for other kinds of cancer.</td>
</tr>
</tbody>
</table>
| The pattern of results for red meat intake is repeated again for processed meat intake. Positive, null, and negative results. Lists of types of cancer that fit in each category. Statistical issues for each finding. | Processed Meat

[Paragraphs omitted: Similar results for processed meat intake]

[Paragraph omitted: Analysis of risk for more specific types of cancer]

| An analysis of each variable—red meat and processed meat—dependent of the other. Statistical methods for this analysis are mentioned, followed by results. | We conducted sensitivity analyses excluding processed meats from the red meat variable to determine whether the risks associated with red and processed meat are independent of each other. . . The positive associations for red meat and cancer of the liver, esophagus, colorectum, and lung all remain . . . Furthermore, the inverse association for red meat and endometrial cancer remained . . . |
|---|
| Remaining interesting statistical details are grouped at the end, to answer any potential concerns the reader may have about the effectiveness of the study. | [Paragraph omitted: Additional statistical analysis showing that results remain significant after controlling for a number of variables.] |

In the following excerpts from the discussion section of the same paper, study results are underlined. Note how the findings mentioned briefly without analysis in the results section are now mentioned again with more analysis and comparison to similar studies. Here, potential limitations of the data are also presented. The results are discussed from strongest proof to weakest. After that, the authors discuss the whole study in general and statistical details and conclude with a clinical application.

<table>
<thead>
<tr>
<th><strong>Comments</strong></th>
<th><strong>Discussion Section</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Topic of the article mentioned first. Note the similarity to the article title. Most significant results also summarized at the beginning of the discussion section. Thus the first paragraph contains no &quot;analysis.&quot;</strong></td>
<td>In this large, prospective investigation of red and processed meat intake in relation to cancer risk, <strong>we found elevated risks for colorectal and lung cancer with both meat types.</strong> Red, but not processed, meat intake was also associated with increased risk for cancer of the esophagus and liver. We observed borderline statistically significant elevated risks for advanced prostate cancer with both red and processed meat intake, for laryngeal cancer with red meat, and for bladder cancer and myeloma and with processed meat intake.</td>
</tr>
</tbody>
</table>

The cancer site most consistently associated with meat intake has been the colorectum. A recent meta-analysis . . . reported elevated risks in the highest category of consumption of . . . meat [9]. **Our study included over 5,000 colorectal cancer cases, and it lends strong support to implicate red and processed meat as risk factors for this malignancy. Consistent with previous studies** [9], we observed a stronger positive association for rectal than colon cancer. |

The strongest finding from the results section is repeated here, then analyzed. Another review of studies is mentioned. The finding agrees with the previous studies. |
Another finding is stated. That this study is the largest of its kind is mentioned. Previous studies are summarized. A potential limitation is mentioned.

The positive associations for both red and processed meat that we report for lung cancer were of similar magnitude to the findings for colorectal cancer. To date, our study includes the largest prospective analysis of meat intake and lung cancer risk. Previous case-control studies have reported elevated risks for lung cancer for those in the highest categories of red meat [17–19], fried red meat [8,19], well-done red meat [17], and processed meat intake [20]. Despite conducting analyses to show that very fine control of smoking history, using a 31-level variable, did not attenuate the lung cancer associations, there remains a potential issue of residual confounding by smoking, because it is such a strong risk factor for this disease.

Another finding is presented. Similar studies are mentioned. That this study is the first one with a prospective methodology is emphasized.

We found a positive association between red meat intake specifically and cancers of the esophagus and liver, and a borderline significant positive association for laryngeal cancer. The first prospective study of meat intake and esophageal cancer was published recently; that study had only 65 cases and found a positive association for processed meat, but not red meat, with esophageal adenocarcinoma [21]. Our study suggests a threshold effect for red meat intake on esophageal cancer risk, beginning at a low level of intake, with no further increase in risk with higher intakes, as reflected in the p-trend (p = 0.13), although it is possible that the referent group had a smaller-than-expected cancer incidence by chance. Data on meat intake and cancers of the liver and larynx are limited, and our study is the first prospective investigation to report on these associations. Two case-control studies reported elevated risks for laryngeal cancer for those in the highest intake categories of red meat intake [22,23] and fried beef/veal [24].

This paragraph contains only findings and comparison to similar studies. There is a brief mention that there are not many studies of this issue.

In our study, those in the highest quintile of processed meat intake had borderline statistically significant elevated risks for myeloma, a malignancy that has not been well-studied for dietary associations, and bladder cancer. A study of two prospective cohorts combined [25], and one case-control study [26], both found elevated risks of bladder cancer for those in the highest categories of processed meat consumption, but another cohort study found no association [27].

An unexpected finding is highlighted with the word “unexpectedly” right at the beginning. Because it is unexpected, the authors clearly show that they adjusted the statistics to avoid confounding from other possible causes. However, the authors also cite studies that offer the opposite results.

Unexpectedly, we found an inverse association between red meat intake and endometrial cancer; this association was not attenuated by adjustment for known risk factors, such as body mass index or menopausal hormone therapy, or by fine control for smoking, which has been inversely associated with this malignancy [28]. Previous studies have reported null [29,30] or positive relations [31] between red meat and endometrial cancer. We also observed inverse associations between processed meat intake and leukemia and melanoma. In contrast to our findings, childhood leukemia has been positively associated with intake of processed meats in a case-control study [32].

There is quite a bit of research on this finding, so the authors group articles and describe them only briefly. They also offer another potential limitation of their findings.

Both red and processed meat intake were positively associated with pancreatic cancer in men, but not women. Red meat has been associated with pancreatic cancer in some [33,34], but not all [35–39] previous cohort studies, as has processed meat in one cohort [34] and several case-control studies [40–44]; although a sex-specific association has not been reported before. Although the association between pancreatic cancer and red or processed meat intake in men was unchanged by fine control for smoking, residual confounding by smoking is still possible.

[Paragraph on other cancers omitted.]
### Previous studies conflict. The authors’ results may explain why.

<table>
<thead>
<tr>
<th>Previous studies of meat intake and prostate cancer are conflicting. Some studies have reported null findings [5,60–66], and others suggest positive associations [67–74]. <strong>Despite finding no association between red or processed meat intake and overall prostate cancer risk, we observed a suggestion of an elevated risk for advanced prostate cancer with both meat types.</strong> If the relation of meat intake to prostate cancer is confined to advanced disease, this could explain some of the inconsistencies in the literature as most previous studies have not specifically addressed advanced prostate cancer.</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Although two recent studies have found a breast cancer/red meat connection, the authors did not find one. They speculate why.</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>With regard to breast cancer, a pooled analysis of eight cohort studies found no association with red meat intake [75]; however, the two most recent prospective studies found positive associations for both red and processed meat [76], specifically for estrogen and progesterone receptor–positive breast cancers in premenopausal women [77]. Although breast cancer risk related to meat intake did not appear to differ by menopausal status in our study, we had very few premenopausal cases (n = 94) and lacked information on hormone receptor status for a large number of cases.</th>
</tr>
</thead>
</table>

[Paragraph on null cancer risks omitted.]

<table>
<thead>
<tr>
<th>No more results given. Interpretation of possible reasons for the results. Reasons are based both on understanding of biological mechanisms and on results of other associational studies.</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Both red meat, regardless of processing procedure, and processed meat can be linked to carcinogenesis by different mechanisms; for example, they are both sources of saturated fat and iron, which have independently been associated with carcinogenesis. Associations between saturated fat and cancer are likely to be related to energy balance in general, whereas iron is thought to contribute to carcinogenesis specifically by generating free radicals and inducing oxidative stress [94]. Most recently, dietary fat was positively associated with breast cancer [95], and iron intake was positively associated with liver [96] and colorectal cancers [97].</th>
</tr>
</thead>
</table>

[Paragraph omitted: More analysis of biological mechanisms for meat/cancer connection.]

<table>
<thead>
<tr>
<th>The authors suggest reasons for the difference in risk for two types of cancer.</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>With regard to the stronger relation of red and processed meat to rectal cancer than to colon cancer, there is variation in several characteristics along the large intestine. . .</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Gap in the research and statistical strengths of study.</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Despite abundant biologic pathways linking meat intake to carcinogenesis at numerous anatomic sites, this is the first comprehensive and prospective analysis. . . A particular strength of this study includes the large size of the cohort. . . An additional strength was that our study provided adequate statistical power to detect associations. Furthermore, recall bias and reverse causation were minimized by. . .</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Limitations of the study and why they are not serious.</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Potential limitations of this study include some degree of measurement error. . . The energy-adjusted correlation coefficients were. . . These correlations compared very favorably to other. . . Although some measurement error remains, the error associated with. . . tends to result in. . . and we attempted to minimize this error by. . . [omitted: additional minimalizations of error and potential errors].</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Conclusion: Main findings repeated again here in the last paragraph. Clinical application of findings.</th>
</tr>
</thead>
</table>

| In conclusion, a diet high in red or processed meat was associated with an elevated risk of both colorectal and lung cancer; in addition, red meat was associated with an elevated risk of esophageal and liver cancer. A decrease in the consumption of red and processed meat could reduce the incidence of cancer at multiple sites. |
4.4.2 Results Structure of an Experimental Study in the IRD(m) Format

Here are some observations about the results structure of an experimental study in the field of immunology. Relative to clinical papers, the entire article is quite long. Although not all basic science journals follow the pattern, this journal uses an IRD(m) format. In other words, the article contains a short introduction, followed by longer results and discussion sections. Then there is a separate methods section at the end in small print. Only when necessary for understanding the results, a few concise summary statements about the method are provided in the results section as well, as detailed below. In this paper, out of 10 pages, 7.5 are devoted to the results and discussion.


The results section of this article is very long and contains several subsections, each with its own subheading. The subheadings denote the main findings:

<table>
<thead>
<tr>
<th>Subheading</th>
<th>Grammatical Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treg cells inhibit priming of TH cells</td>
<td>Sentence with active verb</td>
</tr>
<tr>
<td>Treg cells alter TH cell dynamic activity</td>
<td>Sentence with active verb</td>
</tr>
<tr>
<td>TH cell and Treg cell homing</td>
<td>Nouns: this section is a description</td>
</tr>
<tr>
<td>Lack of stable TH cell–Treg cell interactions</td>
<td>Shorter version of &quot;there are no stable . . . interactions&quot;</td>
</tr>
<tr>
<td>TH cells and Treg cells interact with DCs</td>
<td>Sentence with active verb</td>
</tr>
</tbody>
</table>

Within each subsection, the authors actually describe several cycles of experimentation. In this particular journal, the summary of each cycle is like a mini-research report, containing a hypothesis, description of methods, main result and more detailed results, and an analysis (discussion) of the result. After explaining one procedure’s results, the authors draw a brief conclusion that leads to the next procedure. Note: more often, in other journals it is common not to include the whole cycle. In particular, extra method details and analysis belong in the methods and discussion sections, not in the results section (personal communication, Jeehee Youn).

Treg cells inhibit priming of TH cells

We did initial studies to assess the effect of . . . Treg cells in . . . mice on the priming of . . . TH cells in the pancreatic lymph node. We compared . . . We depleted . . . lymph node cell samples of . . . cells and labeled the cells . . . before transferring them into prediabetic . . . mice. We collected pancreatic and inguinal lymph nodes from the recipient mice 4 d later and measured . . .

In the absence of . . . . . . . TH cells did not proliferate in . . . mice (Fig. 1a) or . . . recipient mice (data not shown), as assessed by . . . [More detailed results.] These results suggested that the presence of endogenous Treg cells in NOD mice suppressed the priming of autoreactive CD4+CD25+ TH cells. To determine whether the differences in the . . . TH cell proliferation in the [two types of] mice were due to the differences in the numbers of . . . Treg cells, we expanded . . . Treg cell populations isolated from . . . donors and used . . . these cells to reconstitute each . . . mouse. Treg cell reconstitution reduced the proliferation of the . . . TH cells . . . [More detailed results.] Thus, the reduced priming and activation of . . . TH cells in these mice was not due to the high frequency of transferred cells. Instead, proliferation of . . . TH in
the pancreatic lymph node inversely correlated with the number of \( T_{reg} \) cells in these mice.

The excerpt above contains two full cycles of hypothesis-methods-results-analysis. The first cycle is the initial experiment with tentative, general findings.

The second cycle asks why this result was found. The experiment supports the new hypothesis. The subsection continues from there with several more cycles of experiments.

The third cycle begins, “Next we determined whether reconstitution of [one type of] \( T_{reg} \) cells was more effective than [another type of] \( T_{reg} \) cells in inhibiting the priming of BDC2.5 T cells in . . . mice.” The third cycle is structured exactly like the first two.

From there, the authors go on to answer more questions about their initial findings to verify that the cause-effect relationship still holds under various conditions. They come to their main conclusion, that \( T_{reg} \) cells inhibit priming of TH cells. All the additional subsections of the results section tell the story of experiments that help the authors describe the cause and effect process in more detail.

Again, for each procedure and its results, there is a cycle in the structure of a mini-journal article. Each cycle contains, at minimum, purpose and results. Some also contain an initial review of the literature and/or research question and a very brief description of the method used. In this journal, most also contain a brief interpretation of the results, but generally, further analysis of the results should be saved for the discussion section.

Although the subject is difficult, the results section is basically a story—the story of the experiments and thought processes of the authors from beginning to end. What makes it easier to read is the repetition of the same content structure and grammatical features in the description of each experimental cycle, as shown in the table below.

### Sections in the Description of an Experimental Cycle

<table>
<thead>
<tr>
<th>Part of Experimental Cycle</th>
<th>Language to Signal the Part</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypothesis in a purpose statement</td>
<td>(In order) to determine . . .</td>
</tr>
<tr>
<td>Methods</td>
<td>We did . . . We compared . . . We depleted . . . We collected . . . etc.</td>
</tr>
<tr>
<td>Results</td>
<td>We detected . . . We almost never found . . . etc.</td>
</tr>
<tr>
<td>Analysis</td>
<td>These results suggested/demonstrated that . . . indicating that . . . Thus . . . etc.</td>
</tr>
</tbody>
</table>

Other features of the authors’ particular style also make it easier to read. The subheadings use a present tense verb, which is shortest and easiest to read, but the main text uses the past tense, which more accurately reflects the fact that the experiments happened in the past. In this particular article, figures and tables are always mentioned in parentheses, never in the sentences of the text. These authors also tend to use the active voice for their main procedures, but passive for procedures that are not as central for understanding the context of the whole paper.

### Example of Less Central Procedure:

In the absence of islet antigen . . . TH cells did not proliferate in . . . mice as assessed by CFSE dilution using flow cytometry.

The authors also make good use of signaling phrases to organize the text.

### Examples:

- In fact, . . .
- Moreover, . . .
- In addition, . . .
- Therefore, . . .
4.5 Showing Certainty about Claims

The results and discussion sections of a research report focus on making claims and then adding support for those claims.

4.5.1 What are “Claims”?

1. Statements about your ideas
2. Statements about your data
3. Statements about other people’s ideas and data

In other words, “claim” is a very general word, and there are many claims in a single journal article.

Here is an example of a claim.

Example:

Basic claim: An increase in smoking among teenagers caused long-term health problems.

When the proof of your idea or data is clear, you should strengthen your claim. When the evidence is less certain, you should limit or weaken your claim. Below are some examples of strengthening and limiting the example above.

Examples of Stronger Claims:

An increase → A sharp increase
caused → undeniably caused, clearly caused, undoubtedly caused, must have caused, etc.
long-term health problems → widespread long-term health problems

You could also add expressions to the beginning of the sentence:

It is clear that an increase . . .
A great deal of evidence leads us to conclude that an increase . . .
We must conclude that an increase . . .

Examples of Limited Claims:

An increase → a probable increase
caused → may have caused, seemed to have caused, contributed to, was one cause of, etc.

Again, you could also add expressions to the beginning of the sentence:

We have reason to believe that an increase . . .
It is possible that an increase . . .

Below is a longer list of expressions from Hyland (2004) that strengthen or limit claims. Note that they have a range of meanings, both positive and negative, so choose carefully after looking at several examples from Google Scholar or other published papers.

4.5.2 Expressions for Strengthening a Claim

**Nouns**
certainty
evidence
the fact that
(without) question

**Adjectives**
assured
certain that
clear
conclusive
is essential
impossible
improbable
inevitable
least
more than
obvious
plain
precise
reliable
sure
ture
unambiguous
undeniable

**Verbs**
conclude
confirm
convince
demonstrate
determine
expect
find
know
it is known that/to
perceive
prove
show
surmise
think

**Adverbs**
will
will not
would not

**Modals**
could not
must

**Adverbs**
precise(ly)
quite
reliable/reliably
sure(ly)
unambiguous(ly)
unarguably
undeniably
undoubted(ly)
unequivocally
unnecessarily
unquestionably

**Interjections**
of course
doubtless
in fact
indeed
no doubt

**Transition**
given that

4.5.3 Expressions for Limiting a Claim

**Adverbs**
admittedly
almost
(not) always
apparently
approximately
basically
conceivable/conceivably
essentially
evidently
formally
frequently
(in) general
generally
hypothetical(ly)
ideally
largely
likely
mainly
maybe
more or less
not necessarily
normally
occasional(ly)
often
ostensibly
partly

**Verbs**
appear
argue
assume

**Adverbs**
believe
claim
deduce
discern
doubt
estimate
guess
hypothesize
imply
indicate
infer
interpret
perceive
postulate
predict
presume
propose
seen (as)
seem
speculate
suggest
suppose
surmise
suspect
tend
Modals
could
may
might
should
should not
would

Nouns
assumption
our belief
certain extent
contention
contention
implication
possibility
prediction
probability
(general) sense
tendency

Adjectives
about
a certain [noun]
around
consistent with
most
open to question
plausible
questionable
uncertain
unclear
unsure

Transition
provided that
4.5.4 Study Limitations

Near the end of the discussion section of your journal article, you should include a paragraph or two addressing the limitations of your study. This is particularly critical in clinical studies, where not acknowledging limitations could lead clinicians to apply your findings before they have been adequately investigated. Here are some examples adapted from Swales of expressions for limiting claims in the discussion section.

**Expressions for limitations of the study:**

- It should be noted that this study has been *primarily* concerned with . . .
- This analysis has *concentrated on* . . .
- The findings of this study are *restricted to* . . .
- This study has addressed *only* the question of . . .
- The *limitations* of this study are *clear* . . .
- We would like to point out that *we have not* . . .

**Expressions for stating conclusions that should NOT be drawn:**

- However, the findings *do not imply*
- The results of this study *cannot be taken as evidence* for . . .
- Unfortunately, we are *unable to determine* from this data . . .
- The *lack of* . . . means that we *cannot be certain* . . .

**Expressions for very limited studies:**

- *Notwithstanding* its *limitations*, this study *does suggest* . . .
- *Despite* its *preliminary* character, the research reported here *would seem to indicate* . . .
- *However exploratory*, this study *may offer some insight* into . . .

4.5.5 Using modals to strengthen or limit a claim

One of the groups of words in each of the two lists above is modals. Modals (can, may, could, etc.) can strengthen or limit a claim. In fact, modals are probably the most common way to show degree of certainty. However, they are very difficult to define for English learners. There is no exact translation among different languages. Here is a more detailed description of the modals. More examples follow later in this chapter.
<table>
<thead>
<tr>
<th><strong>Past and present: no modal. Just a regular verb</strong></th>
<th><strong>Future: WILL</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Description</strong></td>
<td><strong>Examples</strong></td>
</tr>
<tr>
<td>Giving generalizations: statements that people in the field generally agree on</td>
<td>Generalization:</td>
</tr>
<tr>
<td>OR</td>
<td>1. Prevalence of mixed depression, a combination of depression and manic or hypomanic symptoms, <strong>is</strong> high in patients with bipolar disorders.</td>
</tr>
<tr>
<td>Citations: Summarize the findings of others (report, describe, demonstrate, show, prove, etc.)</td>
<td>Benazzi F. Bipolar disorder—focus on bipolar II disorder and mixed depression. Lancet. 2007 Mar 17;369(9565):935-45.</td>
</tr>
<tr>
<td>OR</td>
<td><strong>Citation:</strong></td>
</tr>
<tr>
<td>Reporting certain results: May be proven mathematically. (Rare in biomedical science.)</td>
<td>2. Genetic studies <strong>show</strong> high heritability of the trait, and segregation analysis suggests the presence of an autosomal codominant major gene conferring susceptibility to podoconiosis.</td>
</tr>
<tr>
<td>AND</td>
<td><strong>Prediction:</strong></td>
</tr>
<tr>
<td>Predictions: No or little doubt about the future.</td>
<td>3. Those likely to be sick will face ever increasing premiums, and voluntary coverage <strong>will</strong> continue to decline.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>CAN</strong></th>
<th><strong>Possibility:</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Possibility: It is possible, but will not happen every time.</td>
<td>1. Recent studies <strong>can</strong> result in better outcomes for patients with out-of-hospital ventricular fibrillation.</td>
</tr>
<tr>
<td>OR</td>
<td><strong>Ability:</strong></td>
</tr>
<tr>
<td>Ability</td>
<td>2. How <strong>can</strong> we explain the discrepancy between studies of mite avoidance in children that suggest some benefit [6,8,17] and the data from our study and other studies involving adults that show no improvement in asthma control? [5,10,21]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>WOULD</strong></th>
<th><strong>Limited by a condition:</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Limited by a condition: Often used with an “if” subordinate clause that describes the condition. (The “if” clause is usually unstated when it is clear from the context.)</td>
<td>1. This fundamental restructuring of the payment system <strong>would</strong> achieve both universal coverage and improved efficiency [if . . . ].</td>
</tr>
<tr>
<td></td>
<td>2. [If . . . ] the best impetus for change <strong>would</strong> result</td>
</tr>
</tbody>
</table>
Other uses:

“We would (like to) [reporting verb]. . .”. This is a special expression that allows you to use reporting verbs for yourself.

OR

Past tense of WILL

not from litigation, regulation, or other outside forces, but from within the health care system.


Using a reporting verb with “we”:

3. We would like to emphasize that we cannot prove whether the measured antimyelin antibodies in our patients are antibodies with demyelinating capacity or whether they represent an epiphenomenon of myelin destruction.


4. . . we would recommend the use of a technique including predilation with shorter balloons, the use of longer single stents in order to cover the entire zone of balloon injury. . .


5. Our hypothesis was that high-intensity warfarin would be more effective than moderate-intensity therapy.


SHOULD

Reasonable expectation (more than 50%)

OR

Stating limitations indirectly (a “recommendation” to oneself)

Other Uses:

Making recommendations about future studies or clinical treatment

Reasonable expectation:

1. D-Dimer is a marker of endogenous fibrinolysis and should therefore be detectable in patients with deep-vein thrombosis.


Stating limitations indirectly:

2. Potential limitations of our study should be acknowledged.

Recommendation about future studies:

3. Further analysis of these mice should more clearly define the contribution of SDF1 in this setting and, more globally, to the nonredundant roles for RBP2 demethylase activity in vivo.

**Recommendations about clinical treatment:**

4. … the decision to perform ablation should also take into account the risk of a fatal complication.


5. Both generalists and medical subspecialists should recommend influenza vaccinations to their elderly and high-risk patients.


<table>
<thead>
<tr>
<th>MAY</th>
<th>Possibility (some doubt): Very common for reporting results cautiously.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Possibility: 1. The cohort study had a small number of participants, unaccounted crossover between the groups, and large loss to follow-up, which may have affected the validity of the results.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>COULD</th>
<th>Possibility (more doubt): More cautious than CAN</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Possibility: 1. It is possible that the presence of these mutant p53 proteins in human tumors could negatively affect the outcome of functional p53 restoration depending on how p53 function is restored.</td>
</tr>
<tr>
<td></td>
<td>With “whether”: 2. It remains to be determined whether mutations in MC4R could be one cause of long-term treatment failure.</td>
</tr>
<tr>
<td></td>
<td>Other Uses: Common with “whether”</td>
</tr>
<tr>
<td></td>
<td>Past tense of CAN</td>
</tr>
<tr>
<td></td>
<td>3. Patients were contacted by telephone every 7 to 14 days so that investigators could monitor compliance and safety.</td>
</tr>
<tr>
<td></td>
<td>Past tense of “can”:</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MIGHT</th>
<th>Possibility: Same strength as COULD</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Possibility: 1. Reactive T cells might produce higher levels of interleukin 5, stimulating tissue cosinophilia and subsequent pruritus.</td>
</tr>
</tbody>
</table>
4.5.6 Strength of Claim
Examples in Context

As Turner has noted,

Generally those fields that have fewer variables or variables that can be controlled in the laboratory or tested or simulated mathematically are much more likely to use the present tense to give their conclusions. Those fields... involving human beings or natural process that are hard to isolate in life sciences and medicine are more likely to use model forms (may, etc.) to discuss results.

Thus research reports in biomedical engineering and bioinformatics, for example, will often contain stronger claims than, for example, clinical psychology. The best way to choose the right forms for your own writing is to emulate good writers in your own field. Try searching a collection of PDF files of published articles for modals and the other expressions listed above. Observe how each is used in context, just as the following discussion section is analyzed below.

In the following excerpts from a discussion section, the expressions for strength of claim are underlined. Note that some are exact statistical statements (e.g. significant), and others are more vague (e.g. nearly all). Also note that the authors tend to use different expressions in each part of the discussion section.

The article tested the benefits of providing children under 4 years old with zinc dietary supplements. The first excerpt comes from the very beginning of the discussion section, where the authors summarize the findings they have already presented in the results section. They use a variety of expressions to show the strength or weakness of each claim.


Example:

In our study, zinc supplementation did not result in a significant reduction in overall mortality in children aged 1–48 months in a population with high malaria transmission. However, there was a suggestion that the effect varied by age, with no effect on mortality in infants, and a marginally significant 18% reduction of mortality in children 12–48 months of age (p=0.045). This effect was mainly a consequence of fewer deaths from malaria and other infections. Any effect on mortality in this trial was in addition to a possible effect of vitamin A supplementation... .

The second excerpt suggests several possible interpretations of one result, that zinc supplements did not have a measurable affect on infants less than 1 year old. Note that the authors use “might” and “could” frequently and alternate the two expressions for variety. The discussion is framed at the beginning and end with two other expressions: possible and suggest(ion).
Example:
There are several possible explanations for the absence of effects of zinc supplementation in children younger than 12 months. Infants might have acquired adequate zinc in utero . . . Alternatively, the absence of effect in this age group might be related to the low 5 mg dose used . . . Effects of zinc might be mediated through improvement in immunity . . . and this effect could be restricted in infants . . . Variation in response to zinc supplements in infants in different populations might be expected. Our findings of no effect in infants need further investigation . . . because they could have important implications for targeting of children who would benefit from additional zinc. . . Nutritional and immunological differences might affect responses to infections and survival . . . Thus, the results of this large community-based placebo controlled zinc supplementation trial suggest that . . . zinc supplementation did not have any effect on mortality in infants, but there was a suggestion of reduced mortality in children older than 1 year.

The last section of the discussion offers suggestions for future research. Note the use of “would” in the suggested hypothesis statement. This is a rewritten version of the Yes/No question: “Would a higher dose have a different effect?” As usual, “would” is combined with an implied “if.” “If we did another study, would a higher dose . . . ?” For more information on how to write research questions, see the Introductions chapter.

Example:
Feasible and sustainable methods of enhancing the bioavailable intake of dietary zinc need assessment. We also need to know whether a higher dose would have a different effect in infants, and to elucidate the mechanisms of the effects of zinc and any differences between boys and girls. Our results suggest a need for meta-analysis of all available studies both for mortality and morbidity to make evidence-based recommendation for public health policy to improve mortality, morbidity, growth, and development.


4.6 Recommended Reading

See Chapter 4 of the textbook Academic Writing for Graduate Students for more detailed advice on modifying the strength of claims. Although the book is directed at graduate students, there are also a number of grammar and style tips on other topics that would be helpful even to those who have already earned their degrees.


4.7 Checklists for Evaluating Your Writing

The following two checklists are excerpted from Adam Turner’s English Solutions for Engineering Research Writing (2006). Although they were originally written for evaluating engineering journal articles, all the points are relevant to biomedical research writing.

4.7.1 Results Section Checklist

1. I do not merely describe all of the results, but interpret the important results for the reader. I use words like “significant, moderate, unexpectedly, surprisingly and interestingly,” to interpret the results and not just give a list of results.
2. If appropriate, I have pointed out any problems or inconsistencies with the data (not the same as limitations of the paper).
3. If my results are statistical, I have done all the necessary tests to determine the validity of the results.
4. If my paper does not have a separate “Discussion” section, I have included references that compare my findings with the results in previous research papers.
6. I have used the past tense to talk about the specific results of my paper but I have used the present tense to talk about descriptions of figures or tables and generalizations based on my results of general statements about my whole field.

7. My tables have titles on the top but my figures have captions on the bottom.

4.7.2 Discussion/Conclusion Section Checklist

1. I discuss only the most significant findings and do not simply repeat the results section with more commentary.
2. I have noted any problems with the methods or data. I note the implications of these problems and how they might affect the validity of my conclusions.
3. My discussion section includes references from other papers to either support or compare my research.
4. I have explained why my results differ from previous research if applicable.
5. I have analyzed the structure of papers in my field to understand the relationship between the results, discussion and conclusion sections.
6. I have identified and clearly explained the importance of the findings for the field as a whole.
7. I have mentioned whether my results support or differ from previous research in the field. If they differ, I have attempted to explain why.
8. I have mentioned some possible areas for further research, the importance of the findings or the implications and possible applications of the research (not all are required in all fields).