

Langkah-Langkah Publikasi Jurnal Internasional

Ahmad Hamim Sadewa

Keuntungan Publikasi Ilmiah

1. Komunikasi antar peneliti
2. Sharing idea
3. Prevent duplication
4. Responsibility penerima hibah riset
5. Recognisi pribadi dan institusi
6. Pengembangan penelitian
7. Naik pangkat
8. dan lain-lain

Peran Institusi dalam Publikasi Internasional

1. Membuat kebijakan publikasi untuk setiap dosen (misalnya 1 publikasi/tahun)
2. Kewajiban publikasi bagi penerima hibah penelitian
3. Alokasi dana untuk penelitian, insentif publikasi, membayar biaya publikasi
4. Pelatihan dan pendampingan penulisan manskrip
5. Menciptakan Budaya Menulis, tidak hanya dosen tetapi mahasiswa juga wajib publikasi

Langkah-langkah Publikasi

1. Menyusun manuskrip
2. Menentukan target jurnal
3. Submit manuskrip
4. Reply dari Editor
5. Revisi manuskrip
6. Submit manuskrip yang direvisi
7. Diterima / Ditolak
8. Proof manuskrip
9. Memesan print-off, bayar biaya publikasi
10. Publikasi paper

Menyusun Manuskrip

Manuskrip ditulis sesuai dengan kaidah ilmiah dan bahasa yang benar.

Simpan manuskrip 2-3 minggu, baca lagi (oleh orang lain) untuk meyakinkan sudah ditulis dengan benar.

Manuskrip disusun dalam bagian yang terpisah : introduction, methods, results and discussion, conclusion, reference, tabel dan gambar.

Tampilkan juga Ethical Clearance dan Informed Consent, Material Transfer Agreement

Menyusun Manuskrip

Mulai dari Results : Hasil apa yang penting, tabel atau gambar apa yang akan ditampilkan.

Setelah itu bagian yang paling sulit : Discussion.

Metode dan Latar Belakang

Judul, Author dan Abstrak

Kepustakaan, Ucapan Terima Kasih dll

Menentukan Target Jurnal

Evaluasi manuskrip : bidang yang tepat apa, seberapa besar impactnya.

Tipe publikasi : letter to editors, short communication, technical notes, **original article**, review, invited paper.

Impact jurnal bisa 0 sampai 40-an.

Kawasan : Indonesia, Asia Tenggara, Asia Pasifik, Eropa-Amerika.

Menentukan Target Jurnal

Jenis-jenis jurnal

1. Jurnal internasional dengan impact factor
2. Jurnal internasional tanpa impact factor
3. Jurnal nasional terakreditasi
4. Jurnal nasional tidak terakreditasi

Pastikan : Jurnal berbayar atau gratis!

Submit Manuskrip

Baca dengan teliti petunjuk untuk contributor

Membuat cover letter / letter to the editor

Hardcopy → printout → dikirim via pos

Online submission : manuscript central, load manuskrip ke dalam box yang sudah disediakan.

Apabila berhasil akan mendapatkan ID number.



CLINICAL GENETICS

An International Journal of Genetics and Molecular Medicine

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Clinical Genetics

An International Journal of Genetics and Molecular Medicine

Edited by:
Michael R. Hayden

ISI Journal Citation Reports® Ranking: 2007: 58/132 (Genetics & Heredity)
Impact Factor: 3.181

Clinical Genetics links research to the clinic, translating advances in our understanding of the molecular basis of genetic disease for the practising clinical geneticist. The journal publishes high quality research papers, short reports, reviews and mini-reviews that connect medical genetics research with clinical practice.

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- Information on related prior or concurrent submissions to other publications.
- A list of at least three referees. The list should not name current or prior collaborators.
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Title Page

The first page of the manuscript should contain, on separate lines, the title of the article and the authors' names and institutions. If the title is longer than forty letters and spaces, a short title, not exceeding that limit, will be required for use in the running head. The first page should also include the full postal address, e-mail address, and telephone and fax numbers of the author to whom communications and proofs should be sent. Please be aware that fax number and e-mail address are crucial to our rapid response process.

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Papers must be clearly divided into the following sections: Abstract (not exceeding 200 words); Key Words; Introduction; Materials and Methods; Results; Discussion; Acknowledgements; References; Legends; and Figures/Tables.

Key Words

Four to nine key words for indexing should be given by the author(s) together with the abstract. They should be placed in alphabetical order

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

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
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Manuscripts with Decisions

Manuscript ID	Manuscript Title	Date Submitted	Date Decided	Status	Actions
PED-00068-2005.R1	C117T variant in the survival motor neuron 1 gene was found in the Japanese population	15-Jul-2005 view receipt	17-Oct-2005	ES: Not Assigned • Accept (20-Oct-2005) <i>Archived on 27-Nov-2007</i> view decision letter	
PED-00068-2005	C117T variant in the survival motor neuron 1 gene was found in the Japanese population	10-Feb-2005 view receipt	26-Apr-2005	ES: Asanuma, Kumiko • Major Revision (26-Apr-2005) • a revision has been submitted <i>Archived on 27-Nov-2007</i> view decision letter	a revision has been submitted (PED-00068-2005.R1)

top

Reply dari Editor

Editor akan mengevaluasi manuskrip dan menentukan keputusan.

Kalau belum ada Reply dalam waktu lama, tidak ada kabar dan akan disubmit ke jurnal lain, manuskrip bisa ditarik dengan mengirimkan surat secara resmi.

Reply dari Editor

1. Rejected : manuskrip tidak sesuai dengan “level” atau “jenis” jurnal yang menjadi target, bahasa kurang baik, data kurang valid

Perbaiki manuskrip, submit ke jurnal lain yang sesuai level dan jenisnya.

Reply dari Editor

2. Accepted : manuskrip langsung diterima, kalau manuskrip memang excellent dan sesuai standar jurnal tersebut.

Jarang terjadi, curious : “jurnalnya baik?”

Atau memang manuskripnya bagus, authornya sudah dikenal kemampuannya

Reply dari Editor

3. Revisi, baik major atau minor. Setelah direvisi manuskrip disubmit lagi dengan batas waktu tertentu. Kalau batas waktu terlewati harus submit lagi dari awal.

Paling sering terjadi, masnuskrip direvisi sesuai dengan saran reviewer.

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Clinical Genetics - Manuscript CGE-00036-2006 Friday, February 3, 2006 6:58 AM

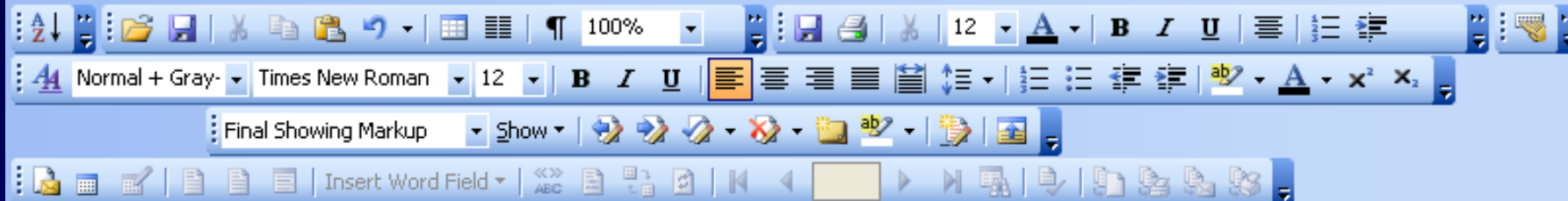
From: "dingen@interchange.ubc.ca" <dingen@interchange.ubc.ca>
To: hamdewa@yahoo.com

Dear Dr. Sadewa:

I am writing to thank you very much for submitting your manuscript entitled "A Novel de Novo Mutation of the *KCNQ2* Gene in Two Japanese Siblings with Benign Familial Neonatal Convulsion." It has now been carefully reviewed by myself and a member of the [Editorial Board](#).

Your manuscript is interesting. However, because of the high increase in recent submissions, we can only publish 40% of the manuscripts we receive. Space constraints dictate that we publish only those manuscripts providing significantly novel insights or new perspectives. Sadly, this means that we are unable to publish many manuscripts of significant quality. Therefore, we have decided not to submit your manuscript to a lengthy [peer review process](#), so as not to delay your submission to another journal.

I appreciate the fact that you have considered Clinical Genetics for the publication of your research. I regret that the decision on this particular manuscript is a negative one, and



Editor-in-Chief

Yukishige Yanagawa

Pediatrics International

These are the editorial comments to the author:

Referee: 1

Comments to the Author

The paper by Sadewa et al., Å novel KCNQ2 mutation and KCNQ3 polymorphism in a Japanese family with BFNC?, describes a family with two siblings affected with BFNC, both carrying a C to T heterozygous substitution in the KCNQ2 gene, predicting a substitution of an arginine by tryptophan, p.R213W, in the S4 segment of the channel protein. Additionally, a novel sequence variant of KCNQ3, c.1241A>G, was detected in the same family. However, this variation was also found in some Japanese healthy controls, suggesting that the KCNQ2 mutation, p.R213W, led to convulsions in the affected siblings.

Comments:



We look forward to receiving your revision.

Sincerely yours,

Yukishige Yanagawa
Editor-in-Chief, Pediatrics International

These are the editorial comments to the author:

Referee 1 Comments:

1) This report describes the occurrence of a silent mutation at codon 28 in exon 2a of survival motor neuron gene (AGC to AGT, C117T variant) not only in control subjects but also SMA patients. The analyzed methods and their results are acceptable and adequate. The authors explained the origin of this C117 variant resulting from a gene conversion mechanism or molecular evolution.

2) I suggest that authors had better to debate about the clinical significance of this C117T mutation in DISCUSSION section. A very recent paper reported the exon 3 is a hot spot for subtle mutation in the SMN1 gene, and it may play a relevant role in SMN protein function (Cusco I et al. Neurology 2004; 63:146-149). Although C117T mutation did not alter the splicing machinery in SMN mRNA, the dosage effect is obscure. The authors should comment on the clinical severity of one SMA patient who had the SMN2 gene with C117T mutation.

Referee 2 Comments:

Revisi Manuskrip

Revisi Minor : menambah diskusi, menambah gambar, koreksi kata, grammar

Revisi Major : menambah data, memperbaiki analisa statistik

Submit Revised Manuskrip

Membuat cover letter.

Menyertakan ID number manuskrip

Manuskrip Diterima

Congratulation! Akan diberitahukan perkiraan tanggal publikasi.

Informasikan ke semua co-author.

Cc: [Empty field]

Subject: FW: Pediatrics International - PED-00268-2006.R2

Attach Files

Plain Text

Rich text toolbar with icons for bold, italic, underline, text color, background color, link, unlink, list, and indent.

We are glad to inform you that your article entitled MS No.: PED-00268-2006.R2 Authors: Sadewa, Ahmad Hamim; Sasongko, Teguh Haryo; ., Gunadi; I Myeong; Daikoku, Kazunari; Yamamoto, Akiyo; Yamasaki, Takemi; Tar Shigenori; Matsuo, Masafumi; Nishio, Hisahide A germline mutation of KCNQ2, p.R213W, in a Japanese family with was accepted on February 2, 2007 for publication in Pediatrics International, and will appear in Vol. 50, No. 3, June 2008.

Send Save as a Draft Cancel

Proof Manuskrip

Mengecek ulang manuskrip, apakah ada kesalahan. Perhatian pada semua bagian manuskrip, terutama tabel, gambar dan referensi.

Awas : perubahan setelah proof disetujui akan dikenakan biaya.

Proof sebaiknya dilihat oleh beberapa orang.


Biaya Publikasi dan Print Off

Beberapa jurnal membebankan biaya publikasi ke author, per halaman. Halaman berwarna lebih mahal dibanding hitam-putih.

Print Off jurnal dikenakan biaya; pesan sedikit saja, sisanya foto copy. Tetapi kalau pesan sedikit, harganya menjadi lebih mahal.

Paper dipublikasikan

Akhir perjuangan panjang dan melelahkan,
diakhiri dengan happy ending.

An aerial photograph of a city skyline. In the foreground, there is a large, well-maintained green lawn. Behind the lawn, there are several tall, modern skyscrapers and residential buildings. The sky is clear and blue. The text is overlaid on the image.

It is not
Publish or Perish

but

Publish and Flourish

Thank you very much