



학회지 SCI core 등재, 영향력 지수 및 인용도 상승계획

대한진단검사의학회지 편집위원장
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국내 SCI Expanded 의편협 학술지 (1)

1. Experimental and Molecular Medicine
 - 영문, 대한생화학분자생물학회, 1996년부터 SCIE / 2002년부터 SCI core
2. Journal of Korean Medical Science
 - 영문, 대한의학회, 1999년부터 SCIE / 2005년부터 SCI core
3. Yonsei Medical Journal
 - 영문, 연세의대, 1998
4. Korean Journal of Radiology
 - 영문, 대한영상의학회, 2001
5. Journal of Veterinary Science
 - 영문, 대한수의학회, 2006

국내 SCI Expanded 의편집 학술지 (2)

6. Journal of Korean Neurosurgical Society (영문, 대한신경외과학회, 2008),
7. Journal of Clinical Neurology' (영문, 대한신경과학회, 2008)
8. Annals of Dermatology'(영문, 대한피부과학회, 2008)
9. Korean Journal of Laboratory Medicine
 - 우리말, 대한진단검사의학회, 2008
10. 'Korean Journal of Parasitology' (영문, 대한기생충학회, 2008)
11. 'Korean Journal of Pathology' (우리말, 대한병리학회, 2008)
12. 'Korean Journal of Physiology and Pharmacology'
 - (영문, 대한생리학회와 대한약리학회 공동 발행, 2008)
13. 'Korean Journal of Orthodontics' (우리말, 대한치과교정학회, 2008)



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[Thomson Scientific](#) : [Press Room](#) : The Scientific Business Of Thomson Reuters Launches Expanded Journal Coverage In Web Of Science By Adding 700 Regional Journals

THE SCIENTIFIC BUSINESS OF THOMSON REUTERS LAUNCHES EXPANDED JOURNAL COVERAGE IN WEB OF SCIENCE BY ADDING 700 REGIONAL JOURNALS

Selected Journals Provide Insight to Scientific Issues That Are Unique to Certain Global Regions

Philadelphia , PA - May 27, 2008 - The Scientific business of Thomson Reuters today announced that 700 new regional journals have been added to Web of Science. The newly identified collection contains journals that typically target a regional rather than international audience by approaching subjects from a local perspective or focusing on particular topics of regional interest.

For more than two years, Thomson Reuters has reviewed thousands of regional journals in all areas of science, social science and arts and humanities. Although selection criteria for a regional journal are fundamentally the same as for an international journal, the importance of the regional journal is measured in terms of the specificity of its content rather than in its citation impact.

"*Web of Science* is regarded as the premier choice in the industry because we focus on the quality of the journals we select – not quantity of journals indexed," said Jim Testa, senior director, editorial development and publisher relations at Thomson Reuters. "By expanding the scope of the regional journal selection process, coverage of material that previously was only available on a limited basis through a few international journals is now accessible in far greater depth to the entire *Web of Science* community. The addition of these first 700 journals will provide meaningful insight to scientific issues that are unique to particular regions throughout the world."

All journals added to the *Web of Science* go through a rigorous selection process. To meet stringent criteria for selection, regional journals must be published on time, have English-language bibliographic information (title, abstract, keywords), and cited references must be in the Roman alphabet. The added journals cover the subject areas of Arts and Humanities, Agriculture Biology & Environmental Sciences, Chemistry, Engineering, Computing & Technology, Life Sciences, Physics, Chemistry, and Earth Sciences, and Social and Behavioral Science.

THE SCIENTIFIC BUSINESS OF THOMSON REUTERS LAUNCHES EXPANDED JOURNAL COVERAGE IN **WEB OF SCIENCE** BY ADDING **700 REGIONAL JOURNALS** (1)

- Philadelphia , PA - May 27, 2008 - The Scientific business of Thomson Reuters today announced that 700 new regional journals have been added to Web of Science. The newly identified collection contains journals that typically target a regional rather than international audience by approaching subjects from a local perspective or focusing on particular topics of regional interest.

THE SCIENTIFIC BUSINESS OF THOMSON REUTERS LAUNCHES EXPANDED JOURNAL COVERAGE IN **WEB OF SCIENCE** BY ADDING 700 REGIONAL JOURNALS (2)

- For more than two years, Thomson Reuters has reviewed thousands of regional journals in all areas of science, social science and arts and humanities.
- Although selection criteria for a regional journal are fundamentally the same as for an international journal, the importance of the regional journal is measured in terms of the specificity of its content rather than in its citation impact.

THE SCIENTIFIC BUSINESS OF THOMSON REUTERS LAUNCHES EXPANDED JOURNAL COVERAGE IN WEB OF SCIENCE BY ADDING 700 REGIONAL JOURNALS (3)

- “*Web of Science* is regarded as the premier choice in the industry because we focus on the quality of the journals we select -- not quantity of journals indexed,” said Jim Testa, senior director, editorial development and publisher relations at Thomson Reuters. “By expanding the scope of the regional journal selection process, coverage of material that previously was only available on a limited basis through a few international journals is now accessible in far greater depth to the entire *Web of Science* community. The addition of these first 700 journals will provide meaningful insight to scientific issues that are unique to particular regions throughout the world.”

THE SCIENTIFIC BUSINESS OF THOMSON REUTERS LAUNCHES EXPANDED JOURNAL COVERAGE IN WEB OF SCIENCE BY ADDING 700 REGIONAL JOURNALS (4)

- All journals added to the *Web of Science* go through a rigorous selection process. To meet stringent criteria for selection, regional journals must be **published on time, have English-language bibliographic information (title, abstract, keywords), and cited references must be in the Roman alphabet**. The added journals cover the subject areas of Arts and Humanities, Agriculture Biology & Environmental Sciences, Chemistry, Engineering, Computing & Technology, Life Sciences, Physics, Chemistry, and Earth Sciences, and Social and Behavioral Science.

대한의학학술지편집인협의회 회원학회 학술지 중 Medline 등재 목록

1. Yonsei Medical Journal, 1963년 [SCIE]
2. Korean Journal of Internal Medicine, 1986년
3. Journal of Korean Medical Science, 1986년9월 [SCI core]
4. Korean Journal of Ophthalmology, 1987년6월
5. Uisahak(의사학), 1992년
6. Korean Journal of Parasitology, 1992년12월 [SCIE]
7. Experimental & Molecular Medicine, 1998년3월 [SCIE core]
8. Korean Journal of Radiology, 2000년3월 [SCIE]
9. Journal of Veterinary Science, 2000년6월 [SCIE]
10. Taehan Kan Hakhoe Chi, 2002~2003년
 - Korean Journal of Hepatology, 2004년3월
11. Korean Journal of Gastroenterology, 2003년7월
12. Taehan Kanho Hakhoe chi (대한간호학회지), 2004년
13. Journal of Preventive Medicine and Public Health, 2005년2월
14. Korean Journal of Laboratory Medicine, 2007년2월 [SCIE]



대한진단검사의학회지 SCI등재 역사 (1)

- 대한임상병리학회지 창간 (1981.12월)
 - 발행인 김기홍, 편집인 김춘원
- 조한익 이사장/ 박명희 편집위원장 (1992-1993)
 - 논문 심사의 질적 향상 노력
- 조한익 이사장/ 김의중 편집위원장 (1994)
 - 책임편집위원 제도 시작
- 지현숙 이사장/ 이갑노 이사장/ 송경순 편집위원장 (1995-2000)
 - 대한의학학술지편집위원회 가입, KoreaMed 및 학술진흥재단 등재
 - PubMed 등재 위한 학술지평가 향상 노력
 - 아카데미아 출판사를 통한 인쇄 질 향상
- 권오현 이사장/ 이경원 편집위원장 (2001-2003)
 - 참고문헌 교정인 제도
 - 대한임상병리학회지 → 대한진단검사의학회지 (2002년6월호) 명칭 변경

第 1 卷 第 1 號 1981

大韓臨床病理學會誌

THE KOREAN JOURNAL OF CLINICAL PATHOLOGY

大韓臨床病理學會 發行

Published by

The Korean Society of Clinical Pathologists

대한진단검사의학회지의 前身 대한임상병리학회지 창간호 1981년

창간호의 투고규정

* 投稿規定 *

1. 투고자격은 대한임상병리학회 회원 및 편집위원회에서 인정하는 사람으로 한다.
2. 원고의 종류는 원저, 증례보고, 심포지움 및 편집위원회에서 위촉한 증설과 조사보고서 등으로 한다.
3. 저재어부 및 순위는 편집위원회에서 결정하고 그내용도 필요에 따라 수정, 보완 또는 삭제할 수 있다.
4. 원고는 국문 또는 영문을 사용하되 국문원고에는 영문초록을 영문원고에는 국문초록을 첨부하여야 한다.
5. 국문원고는 200 자 원고지에 가로쓰기로, 영문원고는 타자기로 이중간격을 두고 타자하여야 한다.
6. 원고내용 표현에서 인명, 지명 등의 고유명사는 원어로, 숫자는 Arabia 숫자로, 도량형은 C. G. S. 단위로 표시하며 약자를 제과 사용할 경우, 처음에 어원을 밝히고 사용한다.
7. 표 (Table) 및 도표 (Figure)의 작성은 흑색으로 하고 제목과 설명은 영문으로 한다.
8. 인용문헌의 배열은 인용된 순서대로 번호에 맞춰야 하고 표기방법은 JAMA 에 준하되 단행본의 경우는 저자명, 제목, 잡지명, 권, 페이지, 연도의 순으로 하고 페이지는 시작되는 페이지만 기재한다. 잡지명의 약호사용시는 약호뒤에 period (.)를 찍지 않는다.

9. 잡지명의 약자는 가급적 Index Medicus에 의거하며 한 단어로 된 잡지명은 약하지 않는다.
 예 1) Post RL, Meritt CR, Albright CD : Membrane adenosine triphosphatase as a participant in the active transport of sodium and potassium in the human erythrocyte. J Biol Chem. 235 : 1796. 1960.
 예 2) Snedecor GW, Cochran GW: Statistical Methods, 6th ed Amer, IOWA State University Press, 1967, p. 45.
10. 현미경 사진은 아트지오텐 또는 특수인쇄로 하는 것을 원칙으로 한다.
11. 기타 언급되지 않은 내용은 일반적인 관례에 준한다.

임원명단

회 장 : 김 기 흥
 부회장 : 김 상 인
 총 무 : 김 춘 원
 재 무 : 조 한 익
 감 사 : 조명준 · 김재식

大韓臨床病理學會誌

The Korean Journal of Clinical Pathology

第1卷第1號(創刊號)

1981年 12月 15日 印刷
 1981年 12月 20日 發行

發行人: 金 箕 洪
 編輯人: 金 春 元

Publisher : Ki Hong Kim, M. D.
 Editor : Choon Won Kim, M. D.

發 行 處 : 大韓臨床病理學會

133 서울特別市 城東區 杏堂洞 17
 漢陽大學校 醫科大學 臨床病理學教室
 293-2111, 3111 # 133, 84

組版·印刷: 大 學 書 林

(非賣品)

대한진단검사의학회지 SCI등재 역사(2)

- 김대원 이사장/ 박찬정 편집위원장 (2004-2005)
 - PubMed/SCI 등재의 Vision과 추진
 - 영문 교정인 제도 (배직현 교수, 서울아산병원 은퇴)
 - 투고규정 정비, 심사위원 수를 3명으로 증가
 - SCI 논문에 대한 대한진단검사의학회지 인용 장려금 마련
 - 온라인 논문투고시스템 개발
 - 저자층 확대를 위해
 - 국외 도서관 발송을 100여곳으로 확대 2005년 4월호부터 보내고 있음

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모 및 변경 내용 최종본 표시(S)

6 4 2 2 4 6 8 10 12 14 16 18 20 22 24 26 28 30 32 34 36 38 40 42 44 46 48 50 52

Abstracts (269 단어)

Background: Human leukocyte antigen (HLA) typing based on polymerase chain reaction (PCR) is rapidly replacing the conventional serological method. This study was intended to evaluate BioSewoom™ HLA-A, -B, -C PCR/SSP Kit (BioSewoom SSP) which had recently been developed in Korea.

Methods: A total of 158 samples from domestic (21) and international (137) HLA proficiency testing (PT) were genotyped with BioSewoom SSP, and its results were compared to consensus results. For comparison with INNO-LiPA HLA-A, -B, -C Typing Kit (INNO-LiPA, Innogenetics, Belgium), 20 samples of Koreans for each HLA-A, -B, -C locus were genotyped with both kits.

Results: Among the 21 samples of domestic PT, BioSewoom SSP showed ambiguities as follows: 4 samples (19.0%) in HLA-A, 6 (28.6%) in HLA-B, and 1 (4.8%) in HLA-C. The ambiguities could be resolved by considering the allele distribution of Koreans. Among the 137 samples from international PT, BioSewoom SSP also showed ambiguities as follows: 12 samples (8.8%) in HLA-A, 26 (19.0%) in HLA-B and 6 (4.4%) in HLA-C. Considering the

삭제됨: 334...296

삭제됨: h...DNA techniques for the typing of human leukocyte antigen (HLA). PCR-reverse sequence specific oligonucleotide probe (reverse-SSOP) and PCR-sequence specific primer (SSP) methods are the most widely used common in routine HLA laboratories. wasis recently

삭제됨: samples... samplesBioSewoom™ SSP Kitthe...the ... of proficiency tests. ... utilizing reverse-SSOP technique,

삭제됨: proficiency testingThere was ...ous cases were found by BioSewoom SSP as many as typing...no discrepancy (무엇과 무엇 사이에 ?)with the consensus in the 21 samples of domestic proficiency testing. However(? 본문에는 domestic proficeincv에도



- 투고자
- 심사위원
- 책임편집위원
- 편집이사

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홈페이지 회원가입

아이디/비밀번호 찾기

대한진단검사의학회지 SCI등재 역사(3)

- 차영주 이사장/ 김정호 편집위원장 (2006-현재)
 - 취임 초부터 SCI 등재 추진
 - 투고규정 재정비 및 영문 번역
 - 비학회원들(외국인 포함) 원고 수용 정책 결정
 - 타 학회 회원 원고 투고 수용
 - 국내외 PhD 급 또는 Non-MD 원고 수용
 - 영문홈페이지 개발
 - 온라인 논문투고시스템 정비
 - 영문학술지 창간 비용 예산 확보 (초기 비용 2100만원)
 - Key title 을 ‘대한진단검사의학회지’에서 ‘Korean Journal of Laboratory Medicine’으로 (2006년11월)
 - Medline 등재신청(2007년4월)
 - DOI/CrossRef 가입
 - Medline 등재허가 통보 받다(2007년11월): 2006년 논문부터
 - SCIE 등재 통보 받다 (2008년4월): 2007년 논문부터



- [Korean Journal of Internal Medicine](#) (Korean J Intern Med, 1226-3303)
Began publication in 1986. Published Irregularly.
- [Korean Journal of Laboratory Medicine](#) (Korean J Lab Med, 1598-6535)
Title changed from Korean Journal of Clinical Pathology in June 2002.
[2002](#) [Jun](#) [Aug](#) [Oct](#) [Dec](#)
[2003](#) [Feb](#) [Apr](#) [Jun](#) [Aug](#) [Oct](#) [Dec](#)
[2004](#) [Feb](#) [Apr](#) [Jun](#) [Aug](#) [Oct](#) [Dec](#)
[2005](#) [Feb](#) [Apr](#) [Jun](#) [Aug](#) [Oct](#) [Dec](#)
[2006](#) [Feb](#) [Apr](#) [Jun](#) [Aug](#) [Oct](#) [Dec](#)
[2007](#) [Feb](#) [Apr](#) [Jun](#) [Aug](#) [Oct](#)
- [Korean Journal of Legal Medicine](#) (Korean J Leg Med, 1225-0589)
Began publication in 1977.
- [Korean Journal of Medical Education](#) (Korean J Med Educ, 1227-8067)
Began publication in 1989.

대한임상병리학회지에서 대한진단검사의학회지로 명칭변경
(2002년6월부터)



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- 2007년 5월 월례집담회
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KSLM Submission

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일반사항

1. 원고는 국문 혹은 영문으로 작성할 수 있으며, 원고의 종류는 원저, 증례 보고, 편집위원회에서 위촉한 증설과 조사보고서 등으로 한다. 증례보고는 국내에서 3회 이상 보고되지 않은 경우를 보고함을 원칙으로 한다. 원고는 또한 다음의 8개 분야로 구분되며, 저자는 자신의 원고에 해당분야구분을 명시하도록 한다. 단, 책임편집위원이 적절한 해당분야가 아니라고 판단하면 분야를 재지정할 수 있다.

2. 윤리성: 사람을 대상으로 연구한 논문인 경우에 헬싱키선언(1964년 발표, 2004년 개정, www.wma.net/e/policy/b3.htm)에 합당하게 연구를 수행하며, 기관의 윤리위원회 또는 임상시험심사위원회(IRB)의 승인을 받고, 필요한 경우에 연구대상자의 동의서를 받았음을 명시해야 한다. 동물실험의 경우, 실험동물의 사육과 사용 등 실험과정, 실험이 행하여진 기관의 윤리위원회 승인을 받았거나, 해당 연구기관의 윤리위원회 규정 또는 NIH Guide for the Care and Use of Laboratory Animals (1996, ILAR [Institute of Laboratory Animal Resources] Committee on NRC, National Academic Press pp125, www.nap.edu/readin-groom/books/labrats/index.html)에 저촉되지 않았음을 기술하여야 한다. 저자들은 논문작성에 사용한 실험자료 원본을 논문출간 시점으로부터 적어도 1년간 보관하고 있어야 하고, 편집위원회의 요청이 있는 경우에 이를 제시하여야 한다.

3. 저작권: 원칙적으로 타 지에 이미 게재된 같은 내용의 원고는 게재하지 않으며, 본지에 게재된 것은 타 지에 게재할 수 없다. 단, 독자층이 다른 타 언어로 된 학술지에 게재하기 위한 경우 등 중복 출판은 양측 편집장의 허락을 받고, 중복출판 원고표지에 각주로 표시하는 등, 다음 문헌에서 규정한 요건을 갖춘 경우에만 가능하다(Ann Intern Med 1997;126:36-47). 원고의 저자들은 모두 논문내용에 대해 의미 있는 기여를 했고, 책임을 지며, 게재승인으로 저작권이 대한진단검사의학회지에 이양되는 내용을 포함한 동의서에 자필 서명하여야 한다.

4. 이해관계 명시(Disclosure of conflict of interest): 연구에 소요된 연구비 수혜내용은 표지하단에 필히 기입하여야 하며, 연구에 관계된, 자료료, 주식 등 이해 관계가 있는 모든 것은 논문표지하단이나 감사의 글 등에 밝혀져야 하고, 이를 모두 명시했음을 원고의 저자 전원의 자필서명이 있어야 한다.

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- 진단유전학(Diagnostic Genetics)
- 검사정보학(Laboratory Informatics)
- 기타 진단검사의학(General Laboratory Medicine)

논문의 작성 요령

1. 원고는 마이크로소프트(MS) 워드 사용을 원칙으로 하며 A4용지에 이중 간격으로 작성한다. Table과 Figure를 제외하고 논문의 양은 A4용지 15매 이내를 원칙으로 한다. 단, 증례보고의 경우는 A4용지 10매 이내로 한다.

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"1, 2, 3"으로 구분한다. 저자들의 성명 뒤에는 학위명이나 자격 등을 명시하되 2가지 이상인 경우에는 대표적인 것 한 가지만 쓴다(예, M.D., Ph.D., M.S., M.T., 등). 국문제목이 30자가 넘거나 영문제목이 40자 넘을 때에는(자간 간격 포함) 줄바꿈을 하여 줄바꿈을 하지 않는다. 초록의 단어의 이해관계(conflict of interest)를 기술한다.

투고 규정 국제적 수준으로 수정함

표지 바로 뒷장을 논문검사를 위한 목표지도 넣고, 목표지에는 편지, 증례보고, 증설, 조사보고서 등 원고의 종류를 명시하고, 분야 구분, 제목(국문과 영문)을 작성하며, 요약제목이 있는 경우 첨부하고, 저자의 소속 및 성명을 기술하지 않는다.

* 전자접수를 위하여 논문접수 또는 수정본의 접수 시에는 반드시 접수용 논문(겉 표지를 포함한 논문 전체)과 심사용 논문(겉 표지를 제외하고, 본문 중에서도 저자가 누구임을 알 수 있는 내용은 모두 암호화된 논문)으로 작성하여 접수한다. 또한 MS워드 파일인 경우에는 "도구" -> "옵션" -> "보안"으로 가서 "개인정보옵션"의 "저장시 개인정보 포함 안함"(저장시 파일 속성의 개인 정보제거)에 표시하고 확인한 후 저장하여 심사자가 저자가 누구인지 알 수 없도록 해야 한다.

2) 초록(abstract): 국문 및 영문원고에 모두 영문초록을 첨부한다. 영문 초록은 250단어 이내로, Background, Methods, Results, Conclusions의 4항목으로 나눈 규정된 형식으로 작성한다. 각 항목은 한 단락으로 작성한다(Uniform Requirements for Manuscripts Submitted to Biomedical Journals, N Engl J Med, 1997;336:309-15). 단, 증례보고의 경우 4항목으로 나누지 않고 한 단락으로 작성하며, 250단어

저작권 이양 및 이해관계 명시에 대한 동의서

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저자가 더 있는 경우에 이 양식을 그대로 복사하여 사용할.

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Professor of Department of Laboratory Medicine
Yongdon Severance Hospital
Yonsei University College of Medicine
146-92 Dogdok-dong, Gangnam-gu
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South Korea

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Publisher: Seoul : Korean Society for Laboratory Medicine

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- Authors:** Chung HJ, Chi HS, Cho YU, Lee EH, Jang S, Park CJ, Seo EJ.
- Department:** Department of Laboratory Medicine, University of Ulsan, College of Medicine and Asan Medical Center, Seoul, Korea.
- Background:** Although cytoplasmic CD79a (cytCD79a) is a highly lineage-specific marker of B lymphoid cells and plays an important role in the diagnosis of acute leukemia, its clinical significance is not fully understood. We aimed to investigate the relationship between cytCD79a positivity and survival probability, and to evaluate the prognostic value of cytCD79a expression in AML with t(8;21) (q22;q22). METHODS: A total of 68 cases of AML with t(8;21)(q22;q22) were diagnosed based on conventional morphology, cytochemistry, flow cytometry, and cytogenetic and molecular genetic analysis. Immunohistochemistry of cytCD79a was performed retrospectively. Laboratory and clinical findings were reviewed. RESULTS: Five patients among 68 AML with t(8;21)(q22;q22) revealed cytCD79a positive reaction; scores for myeloid lineage/B-lymphoid lineage were 5/3-3.5. Among the five cytCD79a positive patients, only one patient was a child. Three patients were with refractory AML or relapsed, and two patients died within 10 months. Median survival time of cytCD79a positive group was shorter (8.0 months) than that (61.3 months) of cytCD79a negative group. The survival probability of the cytCD79a expression group was significantly lower than classical AML with t(8;21)(q22;q22) (P=0.0001). CONCLUSIONS: These findings emphasize the necessity of investigating cytCD79a, especially in AML with t(8;21)(q22;q22), for a different clinical prognostic value.
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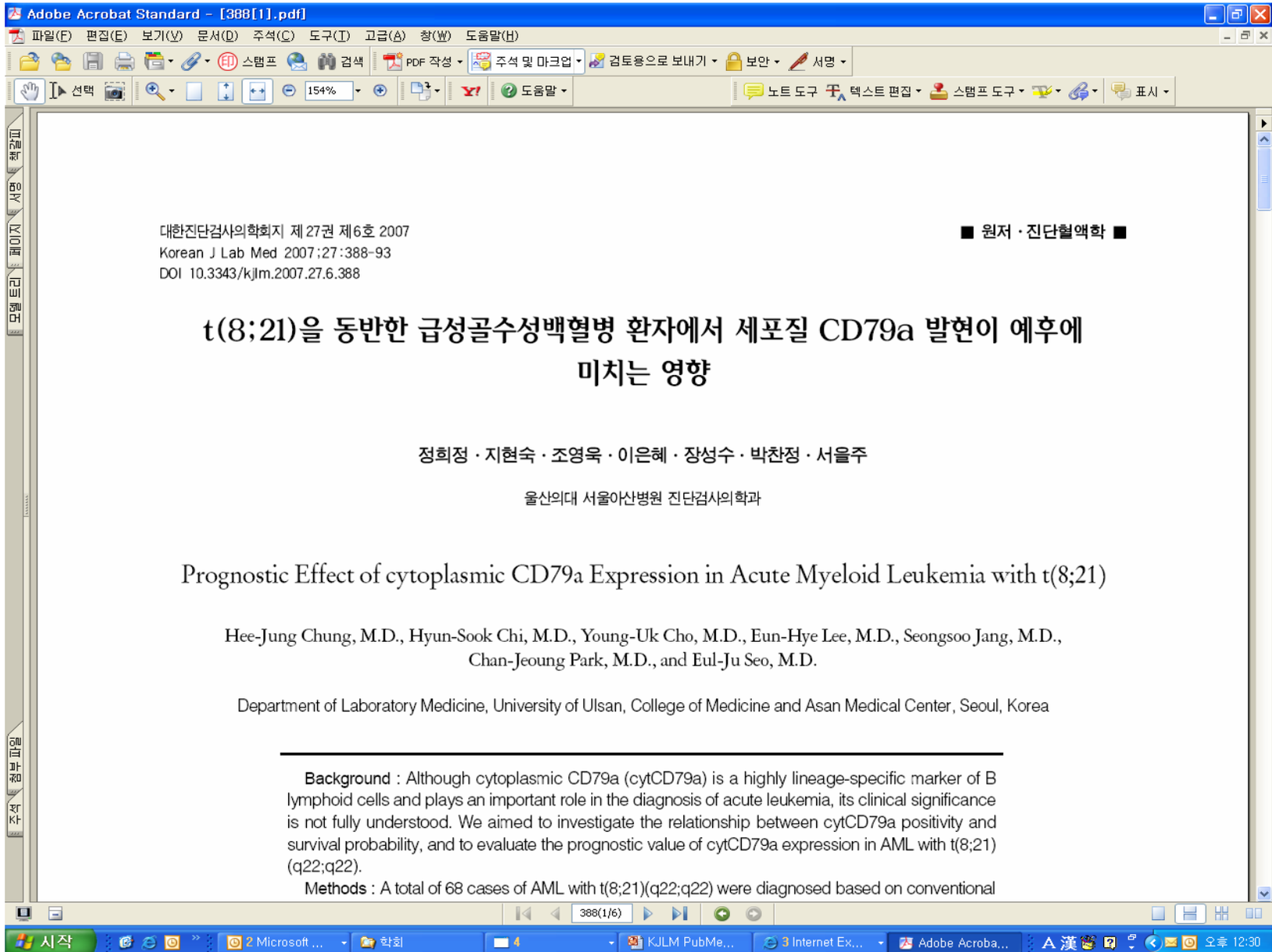
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Institution(s)	Department of Laboratory Medicine, University of Ulsan, College of Medicine and Asan Medical Center, Seoul, Korea
keywords	Acute myeloid leukemia with t(8:21), Cytoplasmic CD79a, Prognosis
Year-vol,-issue	2007 - 27 - 6 Page 388 - 393
Background	Although cytoplasmic CD79a (cytCD79a) is a highly lineage-specific marker of B lymphoid cells and plays an important role in the diagnosis of acute leukemia, its clinical significance is not fully understood. We aimed to investigate the relationship between cytCD79a positivity and survival probability, and to evaluate the prognostic value of cytCD79a expression in AML with t(8:21)(q22;q22).
Methods	A total of 68 cases of AML with t(8:21)(q22;q22) were diagnosed based on conventional morphology, cytochemistry, flow cytometry, and cytogenetic and molecular genetic analysis. Immunohistochemistry of cytCD79a was performed retrospectively. Laboratory and clinical findings were reviewed.
Results	Five patients among 68 AML with t(8:21)(q22;q22) revealed cytCD79a positive reaction; scores for myeloid lineage/B-lymphoid lineage were 5/3-3.5. Among the five cytCD79a positive patients, only one patient was a child. Three patients were with refractory AML or relapsed, and two patients died within 10 months. Median survival time of cytCD79a positive group was shorter (8.0 months) than that (61.3 months) of cytCD79a negative group. The survival probability of the cytCD79a expression group was significantly lower than classical AML with t(8:21)(q22;q22) (P=0.0001).
Conclusion	These findings emphasize the necessity of investigating cytCD79a, especially in AML with t(8:21)(q22;q22), for a different clinical prognostic value. (Korean J Lab Med



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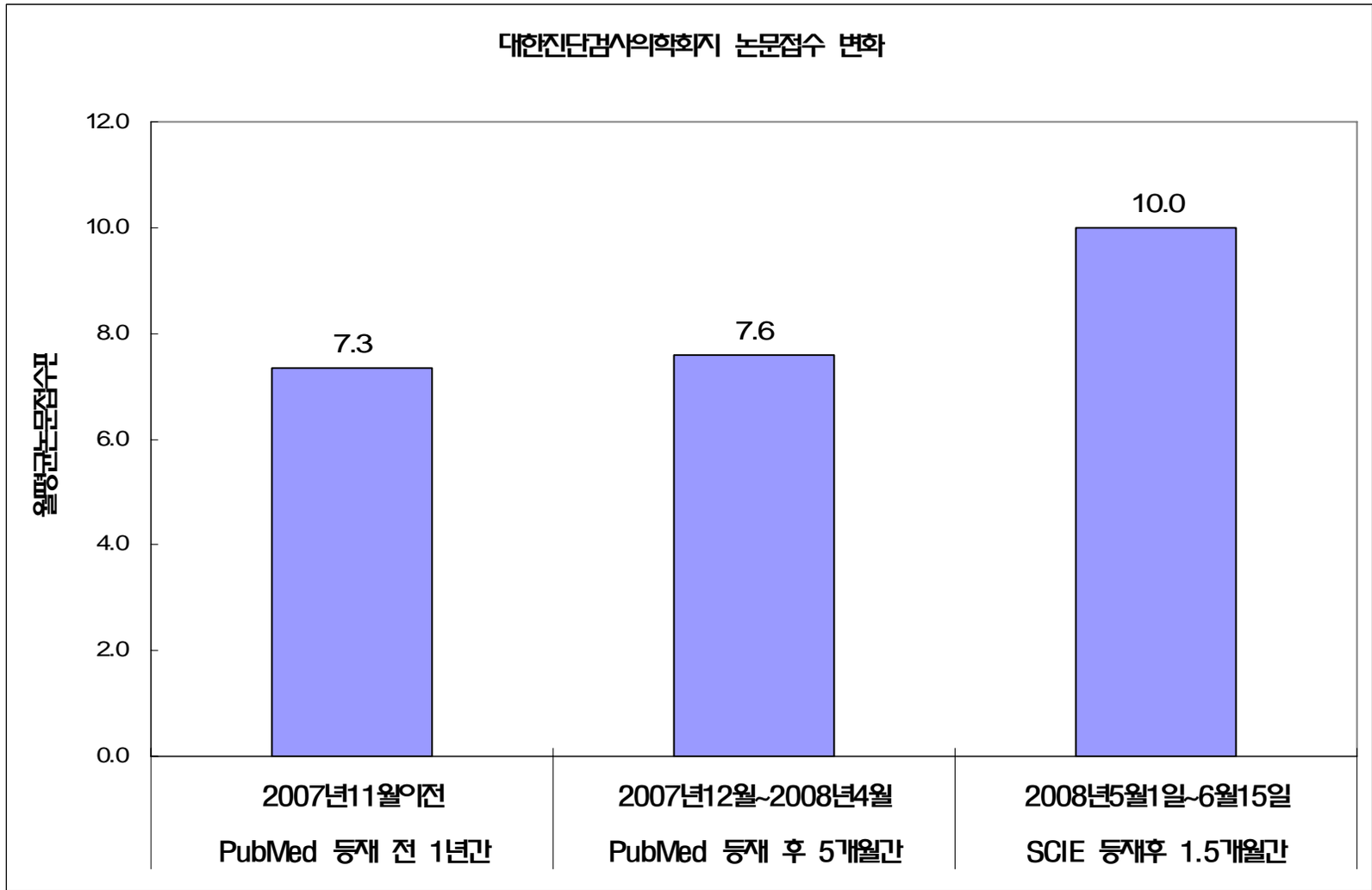
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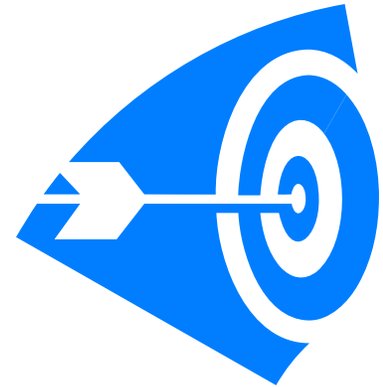


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우리 학회지의 SCI core 진입을 위해

- 우리가 자문할 질문들
 - 우리 학회지의 SCI core 등재가 가능한 것인가?
 - 왜 우리 학회지가 SCI core 등재 되어야 하는가?



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- 사람 → 종업원과 파트너를 개발함으로써 조직의 부가가치를 높여라.
- 문제해결 → 근본의 문제를 지속적으로 해결함으로써 조직 학습을 유도하라. (*Continuously Solving Root Problem Drives Organizationnal Learning.*)
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 - Continuous Improvement (끊임없는 개선, Kaizen)

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2. Editorial Content
3. International Diversity
4. Citation Analysis

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- a) Timeliness of Publication
- b) International Editorial Conventions
- c) English Language Bibliographic Information
- d) Peer Review

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3	CLINICA CHIMICA ACTA	Monthly	SCI core	2.328	2.601	네덜란드
4	THERAPEUTIC DRUG MONITORING	Bimonthly	SCI core	3.032	2.392	미국
5	ADVANCES IN CLINICAL CHEMISTRY	Annual	SCI core	2.44	2.364	미국
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10	CLINICAL CHEMISTRY AND LABORATORY MEDICINE	Monthly	SCI core	1.725	1.741	독일
11	SEMINARS IN DIAGNOSTIC PATHOLOGY	Quarterly	SCI core	0.863	1.667	미국
12	CLINICS IN LABORATORY MEDICINE	Quarterly	SCIE	1.904	1.57	미국
13	CLINICAL LABORATORY	Bimonthly	SCIE	-	1.523	독일
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18	JOURNAL OF CLINICAL LABORATORY ANALYSIS	Bimonthly	SCI core	1.117	0.955	미국
19	JOURNAL OF IMMUNOASSAY & IMMUNOCHEMISTRY	Quarterly	SCI core	0.586	0.614	미국
20	BRITISH JOURNAL OF BIOMEDICAL SCIENCE	Quarterly	SCI core	0.552	0.585	영국
21	PHARMACEUTICAL BIOLOGY	Bimonthly	SCIE	0.397	0.364	미국
22	ANNALES DE BIOLOGIE CLINIQUE	Bimonthly	SCIE	0.342	0.35	프랑스
23	LABMEDICINE	Monthly	SCIE	0.188	0.197	미국
24	ACTA BIOQUIMICA CLINICA LATINOAMERICANA	Quarterly	SCIE	0.125	0.155	아르헨티나
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4	캐나다	1	1
4	네덜란드	1	1
6	프랑스	0	1
6	아르헨티나	0	1
6	크로아티아	0	1
6	대한민국	0	1
	합계	17	27

(4) Citation Analysis

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 - 저자의 피인용 지수 분석 --- 신규 잡지에 중요
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 - 사진, 도표, Table의 인쇄 수준을 높임

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이은엽 (부산의 대)	Kim SY, Kim YJ, Lee SM, Hwang SH, Kim HH, Son HC, Lee EY. Evaluation of the Sysmex UF-100 urine cell analyzer as a screening test to reduce the need for urine cultures for community-acquired urinary tract infection. <i>Am J Clin Pathol.</i> 2007 Dec;128(6):922-5. [Impact factor, 2.939]	Shin JH, Oh YS, Ryang DW. Clinical significance of routine urinalysis. <u><i>Korean J Clin Pathol.</i> 1986;6;33-40.</u>
		Kim CS, Kim KD, Kim DC. Evaluation of usefulness of selective urine culture. <u><i>Korean J Clin Pathol.</i> 1991 May;11(1):109-115.</u>

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Abstract: We report a case of therapy-related acute myeloid leukemia after low-dosed topoisomerase II inhibitor (etoposide) treatment for hemophagocytic lymphohistiocytosis (HLH). A 62-yr-old female patient had previously been treated with a HLH-94 protocol containing a low-dose of etoposide (total dose of 300 mg/m²). Thirty-one months later, the patient was admitted to the hematology department with general weakness and upper respiratory infection symptoms. Peripheral blood smear and bone marrow study revealed acute monocytic leukemia. There was no evidence of myelodysplastic syndrome, and a cytogenetic study showed no chromosomal abnormalities.

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Addresses: Seo, YI (reprint author), Soonchunhyang Univ Hosp, Dept Lab Med, Seoul, South Korea
Soonchunhyang Univ Hosp, Dept Lab Med, Seoul, South Korea
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Hemophagocytic Lymphohistiocytosis and Other Hemophagocytic Disorders

Alexandra H. Filipovich, MD

*Division of Hematology/Oncology, Immunodeficiency and Histiocytosis Program,
Cincinnati Children's Hospital Medical Center, Cincinnati, OH 45229-3039, USA*

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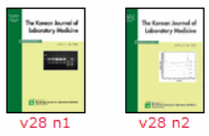
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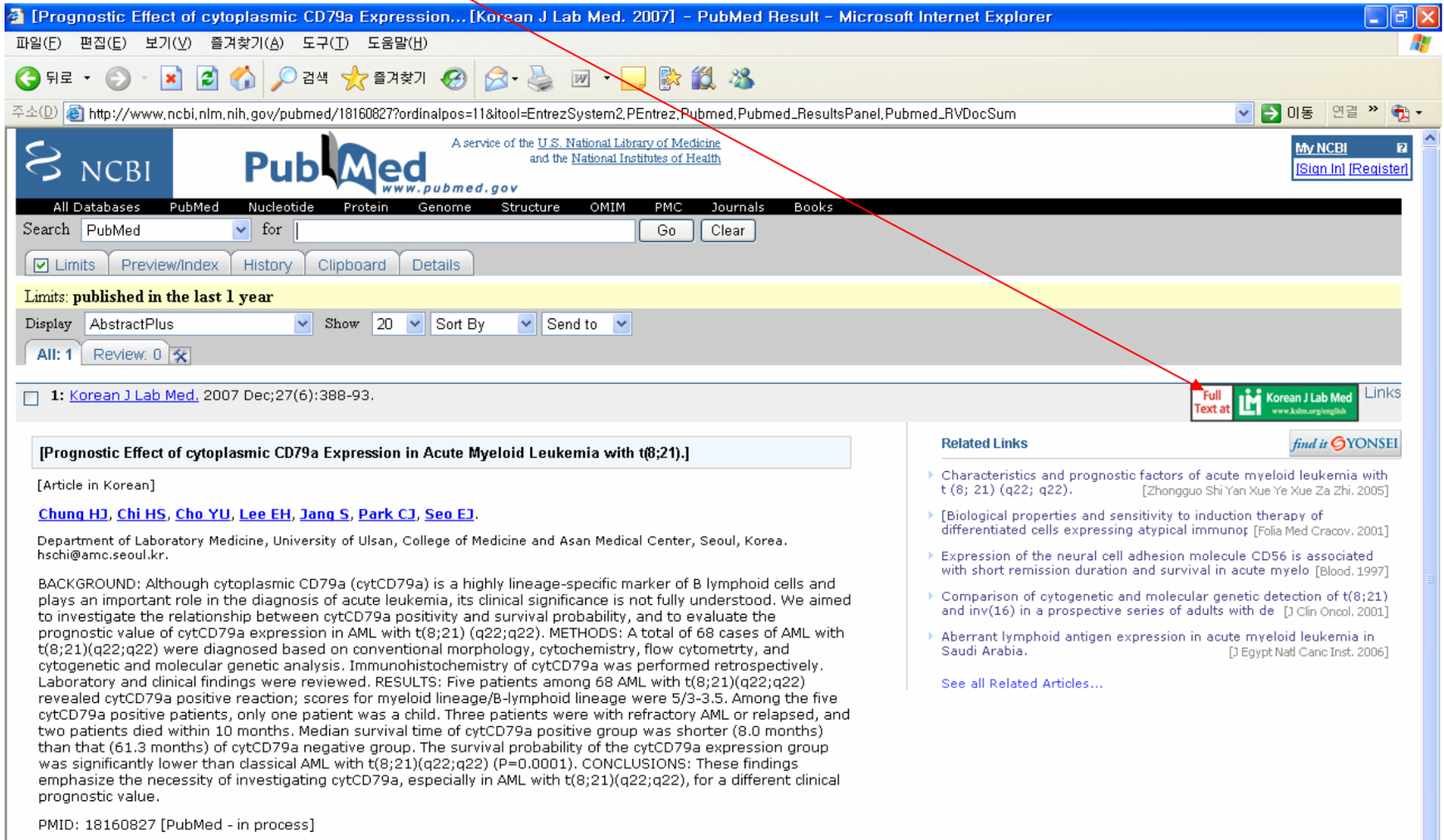
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[Prognostic Effect of cytoplasmic CD79a Expression in Acute Myeloid Leukemia with t(8;21).]

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[Chung HJ](#), [Chi HS](#), [Cho YU](#), [Lee EH](#), [Jang S](#), [Park CJ](#), [Seo EJ](#).

Department of Laboratory Medicine, University of Ulsan, College of Medicine and Asan Medical Center, Seoul, Korea. hschi@amc.seoul.kr.

BACKGROUND: Although cytoplasmic CD79a (cytCD79a) is a highly lineage-specific marker of B lymphoid cells and plays an important role in the diagnosis of acute leukemia, its clinical significance is not fully understood. We aimed to investigate the relationship between cytCD79a positivity and survival probability, and to evaluate the prognostic value of cytCD79a expression in AML with t(8;21) (q22;q22). METHODS: A total of 68 cases of AML with t(8;21)(q22;q22) were diagnosed based on conventional morphology, cytochemistry, flow cytometry, and cytogenetic and molecular genetic analysis. Immunohistochemistry of cytCD79a was performed retrospectively. Laboratory and clinical findings were reviewed. RESULTS: Five patients among 68 AML with t(8;21)(q22;q22) revealed cytCD79a positive reaction; scores for myeloid lineage/B-lymphoid lineage were 5/3-3.5. Among the five cytCD79a positive patients, only one patient was a child. Three patients were with refractory AML or relapsed, and two patients died within 10 months. Median survival time of cytCD79a positive group was shorter (8.0 months) than that (61.3 months) of cytCD79a negative group. The survival probability of the cytCD79a expression group was significantly lower than classical AML with t(8;21)(q22;q22) (P=0.0001). CONCLUSIONS: These findings emphasize the necessity of investigating cytCD79a, especially in AML with t(8;21)(q22;q22), for a different clinical prognostic value.

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Institution(s)	Department of Laboratory Medicine, University of Ulsan, College of Medicine and Asan Medical Center, Seoul, Korea
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Background	Although cytoplasmic CD79a (cytCD79a) is a highly lineage-specific marker of B lymphoid cells and plays an important role in the diagnosis of acute leukemia, its clinical significance is not fully understood. We aimed to investigate the relationship between cytCD79a positivity and survival probability, and to evaluate the prognostic value of cytCD79a expression in AML with t(8:21)(q22;q22).
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Conclusion	These findings emphasize the necessity of investigating cytCD79a, especially in AML with t(8:21)(q22;q22), for a different clinical prognostic value. (Korean J Lab Med

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