The European Register of Specialists in Clinical Chemistry and Laboratory Medicine: Code of Conduct, Version 2 – 2008

Janet McMurray1,*, Simone Zerah2, Michael Hallworth3, Ursula Koeller4, Victor Blaton5, Kamen Tzatchev6, Charis Charilaou7, Jaroslav Racek8, Anders Johnsen9, Karel Tomberg10, Aimo Harmoinen11, Hannsjörg Baum12, Demetrios Rizos13, Janos Kappelmayer14, John O’Mullane15, Giuseppe Nubile16, Silvija Pupure17, Zita Kucinska11e18, Matthias Opp19, Wim Huisman20, Bogdan Solnica21, Henrique Regueno22, Camelia Greigore23, Július Španář24, Greta Strákal25, Josep Queralto26, Hans Wallinder27 and Peter Schuff-Werner28

1 Association for Clinical Biochemistry, London, UK
2 Laboratoire ZTP, Bagnolet, France (SFBC France)
3 Clinical Biochemistry Department, Royal Shrewsbury Hospital, Shrewsbury, UK
4 Institute of Medical Chemistry, Laboratory Diagnostics, Hospital of Lainz, Vienna, Austria
5 Clinical Chemistry Department, AZ Sint Jan Hospital, Brugge, Belgium
6 Clinical Laboratory and Clinical Immunology Department, Medical University, Sofia, Bulgaria
7 Association of Clinical Directors, Biomedical Scientists and Clinical Laboratory Scientists, Nicosia, Cyprus
8 Institute of Clinical Biochemistry and Haematology, Charles University Hospital, Pizen, Czech Republic
9 Clinical Biochemistry Department, National University Hospital, Copenhagen, Denmark
10 North Estonian Regional Hospital, Tallin, Estonia
11 Savonlinnan keskussairaala Laboratorio, Savonlinna, Finland
12 Institut für Laboratoriumsmedizin, Regionale Kliniken Holding Neckar-Schwarzwald GmbH, Ludwigswig, Germany
13 Hormone Laboratory, “Aretaieion” University Hospital, Athens, Greece
14 Clinical Biochemistry and Molecular Pathology Department, University of Debrecen, Debrecen, Hungary
15 Clinical Biochemistry Department, Cork University Hospital, Cork, Ireland
16 Laboratorio Analisi, Ospedale G. Bernabeo di Ortona, Ortona, Italy
17 Clinical Hospital “Gailefers”, Riga, Latvia
18 Department of Physiology, Biochemistry and Laboratory Medicine, Vilnius University, Vilnius, Lithuania
19 Laboratoire National de Santé, Luxembourg, Luxembourg
20 Clinical Laboratory, Medisch Centrum Haaglanden, The Hague, The Netherlands
21 Clinical Biochemistry Department, Jęgiellonien University Medical College, Krakow, Poland
22 Sociedade Portuguesa de Quimica Clinica, Porto, Portugal
23 Laborator Spitalul Clinic de Pediatric Sibiu, University Hospital, Sibiu, Romania
24 Slovak Society of Clinical Biochemistry, Trnava, Slovakia
25 Laboratory Diagnostics Department, General Hospital Murska Sobota, Murska Sobota, Slovenia
26 Servicio de Bioquimica, Hospital Santa Creu I Sant Pau, Barcelona, Spain
27 Clinical Chemistry Department, Aleris Medilab, Taby, Sweden
28 Institute for Clinical Chemistry and Laboratory Medicine, University of Rostock, Rostock, Germany

Abstract

In 1997, the European Communities Confederation of Clinical Chemistry and Laboratory Medicine (EC4) set up a Register for European Specialists in Clinical Chemistry and Laboratory Medicine. The operation of the Register is undertaken by a Register Commission (EC4RC). During the last 10 years, more than 2000 specialists in Clinical Chemistry and Laboratory Medicine have joined the Register. In 2007, EC4 merged with the Federation of European Societies of Clinical Chemistry and Laboratory Medicine (FESCC) to form the European Federation of Clinical Chemistry and Laboratory Medicine (EFCC). A Code of Conduct was adopted in 2003 and a revised and updated version, taking account particularly of the guidelines of the Conseil Européen des Professions Libérales (CEPLIS) of which EFCC is a member, is presented in this article. The revised version was approved by the EC4 Register Commission and by the EFCC Executive Board in Paris on 6 November, 2008.

Keywords: Clinical Chemistry and Laboratory Medicine; Code of Conduct; European Register; registration.


Introduction

Following the expansion of the European Union to 27 countries, and thus the decreased geographical differences between the two organisations, the European Communities Confederation of Clinical Chemistry and Laboratory Medicine (EC4) merged with the Federation of European Societies of Clinical Chemistry and Laboratory Medicine (EFCC) in 2007 to form the European Federation of Clinical Chemistry and Laboratory Medicine (EFCC). EFCC is now the European branch of IFCC. EC4 remains as a sub-section of EFCC with responsibility for the operation of the EC4 Register of Specialists in Clinical Chemistry and Laboratory Medicine.

One of the primary goals of EC4 and EFCC is to stimulate the professional development of the Specialist in Clinical Chemistry and Laboratory Medicine and the maintenance of his\(^1\) professional activities at a very high level (1, 2). To further this aim, in 1997 EC4 set up the European registration system, in which Specialists in Clinical Chemistry and Laboratory Medicine of all countries affiliated to EC4 can be registered (3–5). One of the conditions for registration is that the Specialist in Clinical Chemistry and Laboratory Medicine undertakes to comply with the EC4 Code of Conduct (5). This Code of Conduct is additional to and does not replace any Code of Conduct to which the registrant might be subject in his own country.

The original Code of Conduct was adopted in 2003 (6) and this has now been revised and updated taking account of general changes in the field, and particularly of the statement on common values adopted in 2007 by the Conseil Européen des Professions Libérales (CEPLIS, European Council of the Liberal Professions) (7), of which EFCC is a member.

EC4/EFCC Code of Conduct

General principles

EFCC is the European professional organisation representing Specialists in Clinical Chemistry and Laboratory Medicine, a profession determined by its high level of professional qualifications. The relevant national professional society in each of the EU Member States is represented within EFCC.

In all their work, Specialists in Clinical Chemistry and Laboratory Medicine shall conduct themselves in a manner that does not bring into disrepute the discipline and the profession of Clinical Chemistry and Laboratory Medicine. They shall value integrity, impartiality and respect for persons and evidence and shall seek to establish the highest standards of quality and ethics in their work. Because of their concern for valid evidence, they shall ensure that research is carried out in keeping with the highest standards of scientific integrity. Taking account of their obligations under the law, they shall hold the interest and welfare of patients and those in receipt of their services to be paramount at all times and ensure that the interests of participants in research are safeguarded.

All registrants, having signed an application form, agree to abide by this Code of Conduct. They are also obliged to comply with the Codes of Conduct of their appropriate national registration body and national societies, where appropriate.

Key principles

1. Quality and excellence The Specialist in Clinical Chemistry and Laboratory Medicine shall put his knowledge and ability concerning laboratory diagnostics (including the indication for analyses, the reliability of the results, the interpretation of results and scientific research) at the service of diagnosis, therapy and prevention of human and animal diseases. At all times, he shall act in the best interests of patients, subject to any over-riding legal requirements, with the highest standards of competency and integrity.

2. Continuous professional development In order to optimally fulfil his duties and in accordance with what is regarded as good practice in his profession and having regard to the laws of the country in which he is working, the Specialist in Clinical Chemistry and Laboratory Medicine shall:

- maintain and develop his competence at the highest level of quality by following all relevant (scientific and practical) developments concerning healthcare in general and Clinical Chemistry and Laboratory Medicine in particular, by participating in relevant training courses and other appropriate continuous professional development programmes throughout his working life, and by practising his profession on a regular basis;
- accept assignments only within his area of competence; beyond this limit, he will seek the collaboration of appropriate experts;
- keep up-to-date with statutory codes of practice which affect his work.

The Specialist in Clinical Chemistry and Laboratory Medicine will display his commitment to the profession of Clinical Chemistry and Laboratory Medicine by taking part in the activities of its scientific societies, notably those which promote the profession, and contribute to continuing training of their members.

3. Compliance with codes of ethics and conduct The Specialist in Clinical Chemistry and Laboratory Medicine shall comply not only with the provisions of this Code of Conduct but also with legislation and with any codes of practice and standards relating to his professional work which are applicable in the country in which he is working.

4. Honesty and integrity The professional integrity and intellectual honesty of the Specialist in Clinical Chemistry and Laboratory Medicine shall be the guarantee of his impartiality of analysis, judgment and consequent decisions. The Specialist in Clinical Chemistry and Laboratory Medicine shall at all times avoid deceit in professional

\(^{1}\)Throughout this document he/his are taken for he/she and his/her, respectively.
5. Relationships with others The Specialist in Clinical Chemistry and Laboratory Medicine shall at all times act with courtesy, honesty and integrity in his relationships with patients and others, including professional colleagues, and must not engage in any activity or behaviour which would bring the profession into disrepute or undermine public confidence in the profession.

He must work constructively within a team, and communicate and co-operate with other health professionals and others caring for patients.

He must not abuse his professional position to establish improper relationships with patients, to persuade patients to give or lend money or benefits, to recommend treatments or investigations which are not in the patient’s best interests, or to withhold investigations or treatments.

He must report concerns to employers or regulatory bodies where he believes a colleague’s health, conduct or performance is a threat to a patient.

6. Independence and impartiality The Specialist in Clinical Chemistry and Laboratory Medicine must exercise his professional judgment within the framework of his responsibilities impartially and objectively, after taking into account all relevant circumstances, in the best interests of his patient without pressure from external sources or conflicts of interest. He will ensure that the interests of participants in research are safeguarded and are paramount.

The Specialist in Clinical Chemistry and Laboratory Medicine will serve the individual patient to the best of his ability and provide the general public with such information, within his field of competence, to enable a proper understanding of healthcare matters of public interest.

8. Confidentiality Without prejudice to legislation on privacy applicable in the country where he is working, the Specialist in Clinical Chemistry and Laboratory Medicine will consider himself bound to respect the confidentiality of information obtained by him in his professional work. The Specialist in Clinical Chemistry and Laboratory Medicine will be on his guard against misuse of such information. He will ensure that information about a patient or other individuals is not disclosed to others except in specified circumstances, such as to other health professionals involved in the care of the patient, and, where possible, with the informed consent of the patient.

9. Conflict with moral and ethical beliefs The Specialist in Clinical Chemistry and Laboratory Medicine is not obliged to offer to provide a professional service in ways which conflict with his own moral or religious beliefs, but must respect the moral, religious and cultural beliefs of individual patients. He has an obligation to provide information on where the service requested can most conveniently be obtained from a professional colleague, or details of the institution or professional organisation from which that information can be obtained. If he has agreed to provide a service, he must set aside any personal religious, cultural, philosophical or other convictions. He must ensure equitable access to his services to all who are entitled to use them.

10. Delegation and supervision As head and/or member of the team operating in the Clinical Chemistry and Laboratory Medicine laboratory, the Specialist in Clinical Chemistry and Laboratory Medicine will, given the specific circumstances of the situation concerned:

- obtain a clear definition of the services required of him and/or his team;
- ensure that all activities in the laboratory are organised and executed as accurately and as quickly as possible;
- protect the safety and well-being of his colleagues and be conscious of nature and the environment;
- show respect for superiors, colleagues and subordinates by taking due account of their requirements and aspirations, provided they conform to the laws and ethics of their profession;
- strive for a high level of technical achievement which will also contribute to and promote a healthy and agreeable environment for his colleagues;
- ensure that any member of support staff to whom a task is delegated has the knowledge, skills and competencies necessary to undertake that task effectively and efficiently, and that appropriate supervision is in place;
- retain responsibility for the task delegated, except when the delegatee is at the same level of professional qualification.
11. Professional indemnity insurance The Specialist in Clinical Chemistry and Laboratory Medicine should have in place a form of insurance in respect of potential liabilities to patients and, where applicable, to third parties arising out of his professional work. This should be at a level sufficient to ensure that a justified complainant would be adequately compensated. Such insurance may be provided through a national arrangement for services provided by the state, by an employer, through membership of a professional association or by the individual practitioner. Exceptionally, and by formal prior arrangement, the risk may be borne by the recipient of the service, in Member States where legislation permits such an arrangement. The patient should be made aware of these arrangements.

12. Advertising Specialists in Clinical Chemistry and Laboratory Medicine practise in both the public and private health sectors and the relative distribution between the two varies considerably between the Member States. In Member States where advertising of a Specialist’s services is permitted, any such advertising must be accurate, honest, legal, decent and proportionate, and must focus solely on the professional services offered. It must also conform to any national or EU legislation and guidelines in this area.

Sanctions

Should a Specialist in Clinical Chemistry and Laboratory Medicine not keep to a part of this Code of Conduct, his national regulatory body (where applicable) and his national society will be responsible for determining culpability and sanctions. However, if a registrant is subject to disciplinary sanction (e.g., suspension, removal) from their national register, EC4 will apply the same sanction to the individual in relation to the EC4 Register.

Transparency

The national professional societies are listed, with links, on the EC4 Register website (8) where there are also links to this Code of Conduct and other documents. There are also links to the documents from the website of the European Economic and Social Committee/Single Market Observatory Self- and Co-regulation Database (9). Public access to the names held by the national regulatory bodies is available in some countries. At present, public access to the names of registrants on the EC4 Register is not available but may be in the future. However, this will require consent from each individual.

References