FINAL DRAFT

INTERNATIONAL STANDARD

ISO/FDIS 23191

ISO/TC **249**

Secretariat: SAC

Voting begins on: **2020-04-23**

Voting terminates on: 2020-06-18

Traditional Chinese medicine — Determination of selected Aconitum alkaloids by high-performance liquid chromatography (HPLC)

RECIPIENTS OF THIS DRAFT ARE INVITED TO SUBMIT, WITH THEIR COMMENTS, NOTIFICATION OF ANY RELEVANT PATENT RIGHTS OF WHICH THEY ARE AWARE AND TO PROVIDE SUPPORTING DOCUMENTATION.

IN ADDITION TO THEIR EVALUATION AS BEING ACCEPTABLE FOR INDUSTRIAL, TECHNO-LOGICAL, COMMERCIAL AND USER PURPOSES, DRAFT INTERNATIONAL STANDARDS MAY ON OCCASION HAVE TO BE CONSIDERED IN THE LIGHT OF THEIR POTENTIAL TO BECOME STAN-DARDS TO WHICH REFERENCE MAY BE MADE IN NATIONAL REGULATIONS.



Reference number ISO/FDIS 23191:2020(E)





COPYRIGHT PROTECTED DOCUMENT

© ISO 2020

All rights reserved. Unless otherwise specified, or required in the context of its implementation, no part of this publication may be reproduced or utilized otherwise in any form or by any means, electronic or mechanical, including photocopying, or posting on the internet or an intranet, without prior written permission. Permission can be requested from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office CP 401 • Ch. de Blandonnet 8 CH-1214 Vernier, Geneva Phone: +41 22 749 01 11 Fax: +41 22 749 09 47 Email: copyright@iso.org Website: www.iso.org

Published in Switzerland

Page

Contents

Introduction v 1 Scope 1 2 Normative references 1 3 Terms and definitions 1 4 Abbreviated terms 1 5 Principle 2 6 Reagents 3 7 Apparatus 3 8 Sampling 4 9 Test procedures 4 9.1 General 4 9.2 HPLC-MS procedure 5 9.2.1 Reference solution 5 9.2.3 Chromatographic conditions 5 9.2.4 HPLC-MS procedure 5 9.2.5 Determination 5 9.3.1 Reference solution 5 9.3.2 Test solution 5 9.3.2 Test solution 5 9.3.3 Chromatographic conditions 5 9.3.4 Determination 6 9.3.4 Determination 6	Forew	ord	iv
2 Normative references 1 3 Terms and definitions 1 4 Abbreviated terms 1 5 Principle 2 6 Reagents 3 7 Apparatus 3 8 Sampling 4 9 Test procedures 4 9.1 General 4 9.2 HPLC-MS procedure 5 9.2.3 Chromatographic conditions 5 9.2.4 HPLC-MS conditions 5 9.2.5 Determination 5 9.3.1 Reference solution 5 9.3.2 Test solution 5 9.3.3 Chromatographic conditions 5 9.3.4 Determination 6 9.3.4 Determination 6 9.3.4 Determination 6	Introd	luction	v
3 Terms and definitions 1 4 Abbreviated terms 1 5 Principle 2 6 Reagents 3 7 Apparatus 3 8 Sampling 4 9 Test procedures 4 9.1 General 4 9.2 Test solution 5 9.2.1 Reference solution 5 9.2.2 Test solution 5 9.2.3 Chromatographic conditions 5 9.2.4 HPLC-MS conditions 5 9.3.1 Reference solution 5 9.3.2 Test solution 5 9.3.3 Chromatographic conditions 5 9.3.4 Determination 5 9.3.3 Chromatographic conditions 6 9.3.4 Determination 6 9.3.4 Determination 6	1	Scope	1
4 Abbreviated terms 1 5 Principle 2 6 Reagents 3 7 Apparatus 3 8 Sampling 4 9 Test procedures 4 9.1 General 4 9.2 HPLC-MS procedure 5 9.2.1 Reference solution 5 9.2.2 Test solution 5 9.2.3 Chromatographic conditions 5 9.2.4 HPLC-MS conditions 5 9.2.5 Determination 5 9.3 HPLC-DAD procedure 5 9.3.1 Reference solution 5 9.3.2 Test solution 6 9.3.3 Chromatographic conditions 6 9.3.4 Determination 6 9.3.4 Determination 6	2	Normative references	1
5Principle26Reagents37Apparatus38Sampling49Test procedures49.1General49.2HPLC-MS procedure59.2.1Reference solution59.2.2Test solution59.2.3Chromatographic conditions59.2.4HPLC-MS conditions59.2.5Determination59.31Reference solution59.3.1Reference solution59.3.2Test solution69.3.3Chromatographic conditions69.3.4Determination69.3.4Determination6	3	Terms and definitions	1
6 Reagents 3 7 Apparatus 3 8 Sampling 4 9 Test procedures 4 9.1 General 4 9.2 HPLC-MS procedure 5 9.2.1 Reference solution 5 9.2.2 Test solution 5 9.2.3 Chromatographic conditions 5 9.2.4 HPLC-MS conditions 5 9.2.5 Determination 5 9.3.1 Reference solution 5 9.3.2 Test solution 6 9.3.3 Chromatographic conditions 5 9.3.4 Determination 6 9.3.4 Determination 6	4	Abbreviated terms	1
6 Reagents 3 7 Apparatus 3 8 Sampling 4 9 Test procedures 4 9.1 General 4 9.2 HPLC-MS procedure 5 9.2.1 Reference solution 5 9.2.2 Test solution 5 9.2.3 Chromatographic conditions 5 9.2.4 HPLC-MS conditions 5 9.2.5 Determination 5 9.3.1 Reference solution 5 9.3.2 Test solution 6 9.3.3 Chromatographic conditions 5 9.3.4 Determination 6 9.3.4 Determination 6	5	Principle	2
7 Apparatus 3 8 Sampling 4 9 Test procedures 4 9.1 General 4 9.2 HPLC-MS procedure 5 9.2.1 Reference solution 5 9.2.2 Test solution 5 9.2.3 Chromatographic conditions 5 9.2.4 HPLC-MS conditions 5 9.2.5 Determination 5 9.3.1 Reference solution 5 9.3.2 Test solution 6 9.3.3 Chromatographic conditions 6 9.3.4 Determination 6	6	•	
8 Sampling 4 9 Test procedures 4 9.1 General 4 9.2 HPLC-MS procedure 5 9.2.1 Reference solution 5 9.2.2 Test solution 5 9.2.3 Chromatographic conditions 5 9.2.4 HPLC-MS conditions 5 9.2.5 Determination 5 9.3.1 Reference solution 5 9.3.2 Test solution 5 9.3.3 Chromatographic conditions 5 9.3.4 Determination 6 9.3.4 Determination 6	_	5	
9 Test procedures 4 9.1 General 4 9.2 HPLC-MS procedure 5 9.2.1 Reference solution 5 9.2.2 Test solution 5 9.2.3 Chromatographic conditions 5 9.2.4 HPLC-MS conditions 5 9.2.5 Determination 5 9.3.1 Reference solution 5 9.3.2 Test solution 6 9.3.3 Chromatographic conditions 6 9.3.4 Determination 6 9.3.4 Determination 6	-		
9.2.1Reference solution59.2.2Test solution59.2.3Chromatographic conditions59.2.4HPLC-MS conditions59.2.5Determination59.3HPLC-DAD procedure59.3.1Reference solution59.3.2Test solution69.3.3Chromatographic conditions69.3.4Determination69.3.4Determination6	9	Test procedures 9.1 General	4
9.3 HPLC-DAD procedure 5 9.3.1 Reference solution 5 9.3.2 Test solution 6 9.3.3 Chromatographic conditions 6 9.3.4 Determination 6 10 Test report 6			
10 Test report 6		 9.3 HPLC-DAD procedure 9.3.1 Reference solution 9.3.2 Test solution 9.3.3 Chromatographic conditions 9.3.4 Determination 	
	10	Test report	6
Annex A (informative) Typical HPLC conditions 7	Annex	A (informative) Typical HPLC conditions	7
Annex B (informative) Typical chromatogram of the selected Aconitum alkaloids9	Annex	B (informative) Typical chromatogram of the selected Aconitum alkaloids	9
Annex C (informative) Typical Aconitum herb used in traditional Chinese medicine	Annex	c C (informative) Typical Aconitum herb used in traditional Chinese medicine	
Annex D (informative) Reference values of national and regional limits of Aconitum alkaloids in Aconitum carmichaelii lateral root	Annex		14
Bibliography	Biblio		

ISO/FDIS 23191:2020(E)

Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular, the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation of the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT) see www.iso.org/iso/foreword.html.

This document was prepared by ISO/TC 249, Traditional Chinese medicine.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at www.iso.org/members.html.

Introduction

Aconitum is a genus of about 350 species of flowering plants belonging to the family of Ranunculaceae. The therapeutic use of as many as 76 species of *Aconitum* herbs in traditional Chinese medicine have been documented for a very long time. Among the *Aconitum* herbs, the most well known are processed *Aconitum carmichaelii* lateral root (附子), *Aconitum carmichaelii* root (川乌) and *Aconitum kusnezoffii* root (草乌). Of approximately 80 000 traditional Chinese medicine formulae, around 10,2 % contain *Aconitum* herbs. It is one of the most frequently used groups of herbal medicines in traditional Chinese medicine. Typical *Aconitum* herbs used in traditional Chinese medicine are shown in <u>Annex C</u>, Table C.1.

Aconitum herbs contain *Aconitum* alkaloids which have anti-inflammatory, analgesic and cardiotonic activities. The *Aconitum* alkaloids are a double-edged sword, however. At present, international trade in *Aconitum* products is restricted to a few nations due to the high natural toxicity of crude *Aconitum* products. Unprocessed *Aconitum* is highly toxic. Expert processing is required to reduce (but not eliminate) toxicity, and where individual national regulatory schemes do not ban the herb, it is generally restricted to a high-risk category, such as Schedule 2 in the Hong Kong Chinese Medicine regulations. Also, there are sporadic cases of *Aconitum* alkaloid poisoning due to misuse reported all over the world.

Nonetheless, the toxicity of *Aconitum* herbs can be reduced dramatically with proper processing (such as repeated boiling or steaming), prolonged decoction and dose control. However, testing standards for *Aconitum* alkaloids have not yet been harmonized on an international level, and regulatory authorities in many nations do not adequately differentiate highly toxic forms from less-toxic forms (or even non-toxic forms) of *Aconitum* herbs.

Six kinds of Aconitum alkaloids [aconitine (AC), mesaconitine (MA), hypaconitine (HA), benzoylaconine (BAC), benzoylmesaconine (BMA) and benzoylhypaconine (BHA)] are commonly used as chemical markers for quality control of aconite, determined by the high-performance liquid chromatography (HPLC) method^[1]. The AOAC Official Method 2008.11 also requires the determination of three Aconitum alkaloids, AC, MA and HA, in dietary supplements and raw botanical materials by LC/UV detection with confirmation by LC/MS/MS^[2]. Nevertheless, poisoning cases are still occasionally reported. From 1989 to 2010, 140 cases of *Aconitum* poisoning, including one fatal case, were reported in Hong Kong^[3]. Additionally, 17 cases were reported in Taiwan from 1990 to 1999, 2017 cases in China from 1989 to 2008 and 121 cases in Korea from 1995 to 2007^[4]. Multiple reasons for *Aconitum* poisoning exist and include overdoses, inadequate processing, Aconitum contamination in other herbs, dispensing and management errors and hidden risk factors. In the 17 cases reported in Hong Kong, yunaconitine (YAC), crassicauline A (CCA) and 8-deacetyl-yunaconitine (DYA) were detected instead of AC, MA and HA in the urine samples of the *Aconitum* poisoning patients^[3,4]. As a result, these alkaloids are considered to be hidden risk factors and should be covered in laboratory screenings for toxic compounds^[5]. Therefore, a method to simultaneously determine the levels of these nine alkaloids is needed for quality control of the herb and its products in order to ensure the safe use of these medicinal materials^[6].

This document aims to build a systematic and practical international standard for the determination of *Aconitum* alkaloids with the goal of standardizing the global market, to ensure safe and effective use in clinics and to reduce cases of *Aconitum* alkaloid poisoning.

As national implementation may differ, national standards bodies are invited to modify the limit values of selected *Aconitum* alkaloids in their national standards. Examples of national and regional values are given in <u>Annex D</u>.

HURSI SAMAAND PREMIUM

Traditional Chinese medicine — Determination of selected *Aconitum* alkaloids by high-performance liquid chromatography (HPLC)

1 Scope

This document specifies methods for the determination of the selected *Aconitum* alkaloids, including aconitine, mesaconitine, hypaconitine, benzoylaconine, benzoylmesaconine, benzoylhypaconine, yunaconitine, deacetyl-yunaconitine and crassicauline A.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 3696, Water for analytical laboratory use — Specification and test methods

World Health Organization, Quality control methods for herbal materials, 2011

3 Terms and definitions

No terms and definitions are listed in this document.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at <u>https://www.iso.org/obp</u>
- IEC Electropedia: available at http://www.electropedia.org/

4 Abbreviated terms

For the purposes of this document, the following abbreviated terms apply.

AC	aconitine
BAC	benzoylaconine
BHA	benzoylhypaconine
BMA	benzoylmesaconine
CAS	chemical abstracts service
CCA	crassicauline A
CRS	chemical reference substance
DYA	8-deacetyl-yunaconitine
ESI	electrospray ionization
НА	hypaconitine

ISO/FDIS 23191:2020(E)

HPLC	high-performance liquid chromatography
JAC	jesaconitine
MA	mesaconitine
MRM	multiple reaction monitoring
MS	mass spectrometer
TS	test solution
UV DAD	ultraviolet diode array detector
YAC	yunaconitine

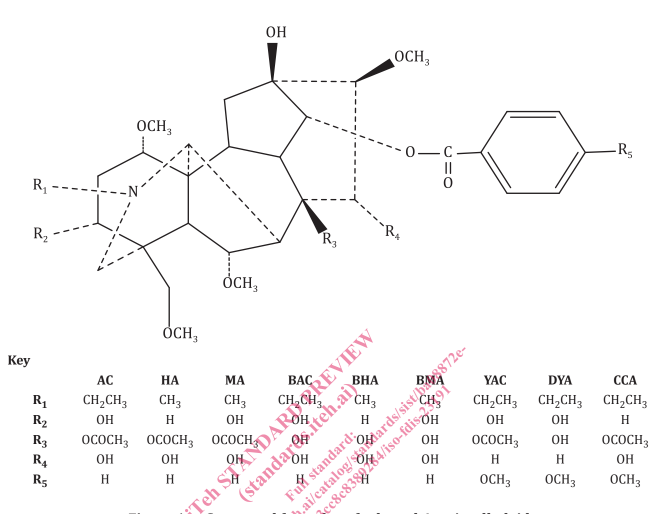
5 Principle

The test solutions (TSs) are analysed by HPLC on a reverse-phase column packed with octadecylsilane bonded silica gel, with 0,1 % formic acid as the mobile phase A, acetonitrile as the mobile phase B and by mass spectrometer (MS) or ultraviolet diode array detector (UV DAD).

Aconitum alkaloids determinable by this method are shown in <u>Table 1</u>. Relevant structural formulae of Aconitum alkaloids are given in <u>Figure 1</u>.

Name	Molecular formula	stanto Stagano.	Molar mass
Name	Molecular Iormula	cull satar 830 no.	g/mol
AC	C ₃₄ H ₄₇ NO ₁₁	ai 200 302-27-2	645,74
MA	C ₃₃ H ₄₅ NO ₁₁	2752-64-9	631,71
HA	C ₃₃ H ₄₅ NO ₁₀	6900-87-4	615,71
BAC	C ₃₂ H ₄₅ NO ₁₀ and At	466-24-0	603,78
BMA	C ₃₁ H ₄₃ NO ₁₀	63238-67-5	589,68
BHA	$C_{31}H_{43}NO_9$	63238-66-4	573,67
YAC	C ₃₅ H ₄₉ NO ₁₁	70578-24-4	659,76
DYA	C ₃₃ H ₅₁ NO ₁₀	110011-77-3	645,79
CCA	C ₃₅ H ₄₉ NO ₁₀	79592-91-9	643,76

Table 1 — Aconitum alkaloids determinable by this method



Structural formulae of selected Aconite alkaloids Figure 1

Reagents 6

9492-4802-92 Silstandar Use only reagents of recognized analytical grade and water conforming with grade 3 as specified in ISO 3696, unless otherwise specified.

- **isopropanol**, C₃H₈O, AR grade;
- dichloromethane, CH₂Cl₂, AR grade;
- acetonitrile, CH₃CN. HPLC grade;
- ethyl acetate, C₄H₈O₂. AR grade;
- methanol, CH₃OH HPLC grade;
- purified water, deionized;
- ammonia TS.

Apparatus 7

The usual laboratory apparatus and the following:

liquid chromatograph, fitted with electrospray ionization (ESI) MS or DAD;

- chromatogaphic column, of stainless steel, 2,1 mm × 100 mm, with 1,7 μm particle size packing of octadecylsilane bonded silica gel;
- ultrasonic bath, power 300 W, frequency 40 kHz;
- **one-mark pipettes**, of capacity 5 ml, 25 ml and 50 ml;
- **conical flasks**, of capacity 100 ml, with glass stopper;
- **vacuum system** (e.g. Büchner flask, Vac-Elut system 1 or peristaltic pump);
- **analytical balance**, capable of weighing to the nearest 0,001 g;
- **microporous membrane**, 0,22 μm, organic phase.

8 Sampling

Sampling shall be carried out in accordance with the method described in the World Health Organization's *Quality control methods for herbal materials, General advice on sampling*. Samples not less than 250 g shall be taken from each batch randomly. Sampling of *Aconitum* genus shall be conducted as follows:

- a) from a batch of five containers or packaging units, take a sample from each;
- b) from a batch of six to 50 units, take a sample from five;
- c) from a batch of over 50 units, sample 10 %, rounding up the number of units to the nearest multiple of 10. For example, a batch of 51 units would be sampled as for 60, i.e. take samples from six packages;
- d) from each container or package selected, take three original samples from the top, middle and bottom of the container or package;
- e) the three original samples shall then be combined into a pooled sample which shall be mixed carefully;
- f) the average sample is obtained by quartering:
 - from the pooled sample, adequately mix into an even and square-shaped heap;
 - divide this diagonally into four equal parts;
 - take two diagonally opposite parts and mix carefully.
 - repeat the process as necessary until the required quantity, to within ± 10 %, is obtained;
- g) using the same quartering procedure, divide the average sample into four final samples, taking care that each portion is representative of the bulk material.

9 Test procedures

9.1 General

The HPLC-MS method applies and the HPLC-DAD method is optional. If the analytical results according to the HPLC-MS method and the HPLC-DAD method are equal, both methods can be applied. If the analytical results are different, the HPLC-MS method shall be used and is preferred for DYA, YAC and CCA because of its high sensitivity. The procedures can be modified if the accuracy and precision are validated. Other similar methods can be applied if they are validated.