

# Cyclopeptide alkaloids

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**Abstract** Cyclopeptide alkaloids are macrocyclic compounds, the ring system of which consists of a hydroxystyrylamine moiety, an amino acid and a β-hydroxy amino acid, and which is substituted with one or two additional units. This review covers scientific literature published since 2006 until today. In the past decade, 39 new cyclopeptide alkaloids have been reported. In addition, absolute or relative configurations of known compounds have been established. New sources of known compounds are listed: Plant families from which cyclopeptide alkaloids have been obtained during the past decade include the Acanthaceae, Malvaceae, Phyllanthaceae and Rubiaceae. Some development concerning total synthesis of cyclopeptide alkaloids are discussed. Finally, a critical overview is given of various biological activities that have been reported, i.e. on the central nervous system (CNS), antimicrobial activity and others.

 $\begin{tabular}{ll} Keywords & Classification \cdot New \\ structures & 2006-2016 \cdot Sources \cdot Total & synthesis \cdot \\ Biological & activities \\ \end{tabular}$ 

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#### Introduction

Cyclopeptide alkaloids are macrocyclic compounds, the ring system of which consists of a hydroxystyrylamine moiety, an amino acid and a  $\beta$ -hydroxy amino acid. This ring is substituted with one or two additional units. Although they occur in various plant families, they are most widely distributed in the Rhamnaceae family, notably the genus Ziziphus (although sometimes the spelling Zizyphus is applied, in this review Ziziphus is used throughout). Over the years they have been covered in various reviews because of their intriguing chemical and biological properties. In 1997 a review on cyclopeptide alkaloids was published covering the literature from January 1985 to December 1995 (Gournelis et al. 1997). About 10 years later, a review on plant cyclopeptides, including cyclopeptide alkaloids, was published covering the literature up to 2005 (Tan and Zhou 2006). El-Seedi et al. (2007) published a review on cyclopeptide alkaloids covering the literature from 1995 to 2005. A chapter was devoted to cyclopeptide alkaloids from higher plants in the book series "The Alkaloids", covering the literature up to 2008, by Morel et al. (2009). Since then, no comprehensive review on the cyclopeptide alkaloids has been published anymore. In view of the limited accessibility of book chapters, the present review was conceived in order to cover new findings and developments since 2005, i.e. covering the literature published from 2006 until now (last SciFinder® access on 27 September 2016).



## Classification

The cyclopeptide alkaloids sensu stricto contain a 13-, 14- or sometimes 15-membered macrocyclic ring (Gournelis et al. 1997; Joullié and Richard 2004). In the 13-membered ring compounds, this macrocyclic ring is closed through an ether bridge in the *m*-position of the styrylamine unit, whereas in the 14-membered ring compounds the ring is closed in the p-position. As exemplified for 13-membered ring compounds in Fig. 1, cyclopeptide alkaloids typically contain four building blocks (A, B, C and D) and in particular series of compounds a fifth building block E is present between A and B. "A" is a basic terminal amino acid, usually with a primary, monomethylated or dimethylated amino group; "B" is a β-hydroxy amino acid; "C" is an amino acid taking part in the macrocyclic ring; "D" is the hydroxystyrylamine unit; and "E" (if present) is an amino acid. In this way the macrocyclic ring contains two amide bonds and one ether linkage. According to the size of the macrocyclic ring (13- or 14-membered) and the number of building blocks (4 or 5), cyclopeptide alkaloids can be classified as belonging to the 4(13), 5(13), 4(14) or 5(14) type. The 14-membered ring class is the largest one, and the 4(14) and 5(14) groups are subdivided according to the nature of the β-hydroxy amino acid (building block B). In this way, the 4(14) group has been subdivided into the

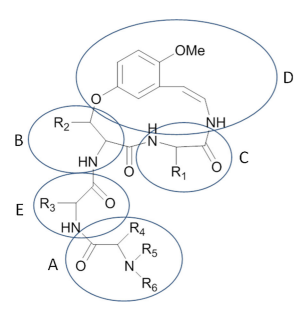
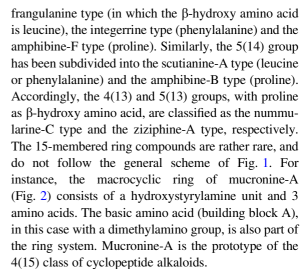


Fig. 1 General structure of 13-membered cyclopeptide alkaloids



The 14-membered cyclopeptide alkaloids have also been categorized according to the macrocyclic ring (El-Seedi et al. 2005), substructure 1 covering the frangulanine, intergerrine and scutianine-A type; substructure 2 covering the pandamine and hymenocardine type (see below); substructure 3 being a unique substructure represented by anorldianine; substructure 4 covering the amphibine-F type with proline; and substructure 5 covering the amphibine-B and -F type alkaloids without proline.

In addition to the cyclopeptide alkaloids sensu stricto, also linear peptide alkaloids, in which the macrocyclic ring system is broken, and neutral compounds, in which the basic amino acid (building block A) is missing, can be distinguished (Gournelis et al. 1997). Some particular features of the neutral cyclopeptide alkaloids are discussed in "Neutral cyclopeptide alkaloids" section. Although Tan and Zhou (2006) have proposed a classification system of plant cyclopeptides

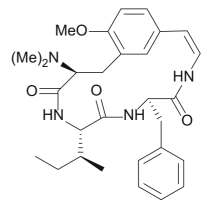


Fig. 2 Mucronine-A, a 15-membered cyclopeptide alkaloid



**Table 1** 4(13) Cyclopeptide alkaloids (nummularine-C type)

No.	Name	Plant origin	Family	Chem. formula	$M_{\rm r}$	References
1	Sativanine-N	Ziziphus sativa Gaertn.	Rhamnaceae	$C_{26}H_{38}N_4O_5$	486	Singh et al. (2006)
2	Sativanine-O			$C_{32}H_{34}N_4O_5$	554	
3	Xylopyrine-A	Ziziphus xylopyrus (Retz.) Willd.	Rhamnaceae	$C_{31}H_{40}N_4O_5$	548	Singh et al. (2007a)
4	Xylopyrine-B			$C_{34}H_{38}N_4O_5$	582	Pandey et al. (2008a)
5	Xylopyrine-D			$C_{30}H_{38}N_4O_5$	534	
6	Xylopyrine-E			$C_{28}H_{34}N_4O_5$	506	
7	Jubanine-E	Ziziphus jujuba Mill.	Rhamnaceae	$C_{33}H_{36}N_4O_5$	568	Pandey et al. (2008a)
7	Joazerine (=Jubanine-E)	Ziziphus joazeiro Mart.	Rhamnaceae	$C_{33}H_{36}N_4O_5$	568	Singh et al. (2012a)

in which the cyclopeptide alkaloids (type I) are subdivided as types Ia1, Ia2, Ia3, Ib and Ic, in this review the classification system described above used in the previous reviews by Gournelis et al. (1997) and El-Seedi et al. (2007) was applied.

### Structures of new compounds

Cyclopeptide alkaloids of the 4(13) type (numularine-C type)

Since 2006, a total of 7 new cyclopeptide alkaloids of the 4(13) type have been reported, more in particular

belonging to the nummularine-C type with proline as the β-hydroxy amino acid (building block B) (Table 1). Sativanine-N and sativanine-O were obtained from stem bark of *Ziziphus sativa* Gaertn (Singh et al. 2006). Sativanine-N (1) contains isoleucine both as ring-bound amino acid (building block C) and as basic terminal amino acid (building block A) with an unsubstituted primary amino group. In sativanine-O (2), 2 phenylalanine units are present instead, still with a primary amino group. From root bark and bark of *Ziziphus xylopyrus* (Retz.) Willd. (although originally referred to a *Zizyphus xylopyra*) 4 cyclopeptide alkaloids of the same type have been reported, all of them containing a phenylalanine unit



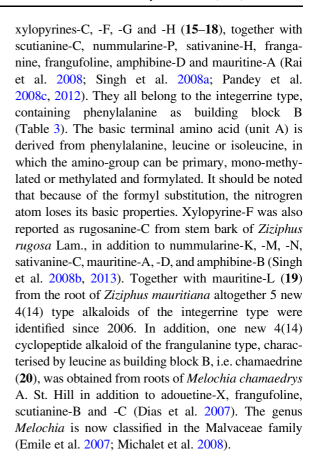
as ring-bound amino acid (building block C). The basic terminal amino acid (building block A) is *N*,*N*-dimethyl-leucine in xylopyrine-A (3), *N*,*N*-dimethyl-phenylalanine in xylopyrine-B (4), *N*,*N*-dimethylvaline in xylopyrine-D (5), and valine in xylopyrine-E (6) (Singh et al. 2007a; Pandey et al. 2008a). A cyclopepeptide related to xylopyrine-B, in which the A-unit is *N*-methyl-phenylalanine, was isolated from bark of *Ziziphus jujuba* Mill. and named jubanine-E (7), together with the known compound nummularine-K (Pandey et al. 2008a, d). The same compound was obtained from *Ziziphus joazeiro* Mart. and named joazerine, in addition to the known compound nummularine-M (Singh et al. 2012a, b).

Cyclopeptide alkaloids of the 5(13) type (ziziphine-A type)

In the same time frame, also 7 new cyclopeptide alkaloids of the 5(13) type, more in particular of the ziziphine-A type, again characterised by proline as the β-hydroxy amino acid (building block B), have been described (Table 2). Sativanine-M isolated from Ziziphus sativa bark contains isoleucine as building block C (ring-bound amino acid), valine as additional building block E, and an alanine derivative as basic terminal amino acid (building block A), i.e. N-formyl, N-methyl-alanine (8). Also nummularine-P was reported (Pandey et al. 2008b). Five jubanines (F-J) (9–13) belonging to the same type have been obtained from roots of Ziziphus jujuba, in addition to the known compounds nummularine-B, daechuine-S3 mucronine-K (Kang et al. 2015). They all have isoleucine or valine as building blocks C or E in various combinations, and N-methyl- or N,Ndimethyl-alanine as basic terminal amino acid. Finally, mauritine-M (14) was reported from the root of Ziziphus mauritiana Lam. (Panseeta et al. 2011). Mauritine-M contains N-methyl-leucine, proline, isoleucine and tryptophane as building blocks A, B, C and E, respectively (D being the hydroxystyrylamine unit).

Cyclopeptide alkaloids of the 4(14) type (integerrine and frangulanine type)

In addition to the xylopyrines discussed above, belonging to the 4(13) class, some other new xylopyrines have been isolated from root bark or bark of the same species (*Ziziphus xylopyrus*), showing a 4(14) structure, i.e.



### Cyclopeptide alkaloids of the 5(14) type

Only one new cyclopeptide alkaloid belonging to the scutianine-A type (with leucine or phenylalanine as the β-hydroxy amino acid unit (building block B) has been reported since 2006, i.e. oxyphylline-A (21) from stem bark or Ziziphus oxyphylla Edgew. (Table 4), together with nummularine-R (Inayat-Ur-Rahman et al. 2007). The terminal basic amino acid (unit A) is N,N-dimethyl-phenylalanine, the additional intermediary amino acid (unit E) proline. In addition to sativanine-K, one 5(14)-type cyclopeptide alkaloid has been isolated from root bark of Ziziphus mauritiana, i.e. mauritine-K (22), in which the  $\beta$ -hydroxy amino acid unit (building block B) is proline (amphibine-B type), the basic terminal amino acid leucine (unit A), the ring-bound amino acid (unit C) as well as the additional amino acid (unit E) leucine (Singh et al. 2007b). Six new cyclopeptide alkaloids (23–28) belonging to the same amphibine-B type have been isolated from stems of Ziziphus apetala Hook. f. in addition to the known compounds mauritine-A and



**Table 2** 5(13) Cyclopeptide alkaloids (ziziphine-A type)

No.	Name	Plant origin	Family	Chem. formula	$M_{\rm r}$	References
8	Sativanine-M	Ziziphus sativa Gaertn.	Rhamnaceae	$C_{30}H_{43}N_5O_7$	585	Pandey et al. (2008b)
9	Jubanine-F	Ziziphus jujuba Mill.	Rhamnaceae	$C_{28}H_{41}N_5O_6$	543	Kang et al. (2015)
10	Jubanine-G			$C_{29}H_{43}N_5O_6$	557	
11	Jubanine-H			$C_{30}H_{45}N_5O_6$	571	
12	Jubanine-I			$C_{30}H_{45}N_5O_6$	571	
13	Jubanine-J			$C_{31}H_{47}N_5O_6$	585	
14	Mauritine-M	Ziziphus mauritiana Lam.	Rhamnaceae	$C_{38}H_{50}N_6O_6$	686	Panseeta et al. (2011)

mauritine-F (Han et al. 2011). Apetaline-A (23) and -B (24) consist of phenylalanine as the β-hydroxy amino acid (building block B) and the ring-bound amino acid (unit C), valine as additional amino acid (unit E), and an unusual terminal unit (A), i.e. 2-(hydroxylimino)-propanoic acid (which can be considered as an alanine derivative) in apetaline-A, and an imidizolidine-4-one structure (formed by bridging one methyl of N,N-dimethyl-alanine and the N-atom of the preceding

valine unit) in apetaline-B. Another new compound was identified as epimauritine-A (25), being the C-34 epimer of mauritine-A (C-34 is the chiral carbon of the terminal alanine moiety). In addition, also epimauritine-A *N*-oxide (26) and mauritine-A *N*-oxide (27) were obtained. Apetaline-C (28) is similar to (epi)mauritine-A, but contains *N*-formyl, *N*-methylamine-alanine as terminal amino acid; the absolute configuration at C-34 could not be determined.



**Table 3** 4(14) Cyclopeptide alkaloids (integerrine- and frangulanine-type)

$$R_2$$
  $R_3$   $R_4$ 

20 Chamaedrine

15 Xylopyrine-C	$R_1 = R_2$	Benzyl	$R_3$	$CH_3$
			$R_4$	Н
16 Xylopyrine-F	$R_1 = R_2$	$CH_2CH(CH_3)_2$	$R_3 = R_4$	Н
<b>17</b> Xylopyrine-G	$R_1$	Benzyl	$R_3$	$CH_3$
	$R_2$	$CH_2CH(CH_3)_2$	$R_4$	СНО
18 Xylopyrine-H	R <sub>1</sub>	Benzyl	$R_3$	$CH_3$
	$R_2$	$CH(CH_3)_2$	$R_4$	Н
19 Mauritine-L	$R_1 = R_2$	CH(CH <sub>3</sub> )CH <sub>2</sub> CH <sub>3</sub>	$R_3$	CH <sub>3</sub>
			$R_4$	Н

(15 - 18: no relative configurations assigned)

No.	Name	Plant origin	Family	Chem. formula	$M_{\rm r}$	References
15	Xylopyrine-C	Ziziphus xylopyrus (Retz.) Willd.	Rhamnaceae	$C_{36}H_{36}N_4O_4$	588	Singh et al. (2008a)
16	Xylopyrine-F			$C_{29}H_{38}N_4O_4$	506	Pandey et al. (2008c)
17	Xylopyrine-G			$C_{34}H_{38}N_4O_5$	582	Pandey et al. (2012)
18	Xylopyrine-H			$C_{32}H_{36}N_4O_4$	540	
16	Rugosanine-C (=xylopyrine-F)	Ziziphus rugosa Lam.	Rhamnaceae	$C_{29}H_{36}N_4O_4$	506	Singh et al. (2013)
19	Mauritine-L	Ziziphus mauritiana Lam.	Rhamnaceae	$C_{30}H_{40}N_4O_4$	520	Panseeta et al. (2011)
20	Chamaedrine	Melochia chamaedrys A. St.Hill.	Malvaceae	$C_{36}H_{41}N_5O_4$	607	Dias et al. (2007)

Recently three new cyclopeptide alkaloids have been obtained from root bark of *Hymenocardia acida* Tul. (Phyllanthaceae) (Tuenter et al. 2016). In addition to the known major compound hymenocardine, also hymenocardine *N*-oxide (29), hymenocardinol (30) and hymenocardine-H (31) were obtained. These cyclopeptide alkaloids do not contain the usual hydroxystyrylamine building block, but rather a *p*-hydroxyphenylethylamine unit: The –CH = CH– moiety is replaced by a –CH(OH)–CH<sub>2</sub>– group in hymenocardinol (30). Hymenocardinol can be considered as the

reduced analogue of hymenocardine, possessing a hydroxyl- instead of a keto-functionality. Whereas hymenocardine and hymenocardinol consist of *N*,*N*-dimethyl-isoleucine, valine, tryptophane and valine as building blocks A, B, C and E, respectively, the new cyclopeptide alkaloid hymenocardine-H consist of *N*,*N*-dimethyl-isoleucine, leucine, histidine and isoleucine as building blocks A, B, C and E, respectively. Because of the presence of histidine, an unusual amino acid in cyclopeptide alkaloids, the name hymenocardine-H (31) was adopted for this compound.



Table 4 5(14) Cyclopeptide alkaloids

1-OH-Benzyl 21 Oxyphylline-A  $R_1$ 

23 Apetaline-A

 $R_1$ Benzyl

 $R_2$ 

 $CH(CH_3)_2$ 

24 Apetaline-B

 $R_1$ Benzyl  $R_2$  $CH(CH_3)_2$ 

29 Hymenocardine N-oxide

 $X_1$ C=O N<sup>+</sup>-O

30 Hymenocardinol

 $X_1$ CH-OH

22 Mauritine-K

 $R_1 = R_2 = R_5$ CH(CH<sub>3</sub>)CH<sub>2</sub>CH<sub>3</sub>  $R_3 = R_4$ 

(22: no relative configurations assigned)

 $R_1$ Benzyl 25 Epimauritine-A  $R_2$ CH(CH<sub>3</sub>)<sub>2</sub> 26 Epimauritine-A N-oxide 27 Mauritine-A N-oxide  $R_3 = R_4 = R_5$ 

28 Apetaline-C

Benzyl CH(CH<sub>3</sub>)<sub>2</sub>

 $R_3 = R_5$  $CH_3$ 

СНО

31 Hymenocardine-H

No.	Name	Plant origin	Family	Chem. formula	$M_{\rm r}$	References
21	Oxyphylline-A	Ziziphus oxyphylla Edgew.	Rhamnaceae	$C_{42}H_{45}N_5O_6$	715	Inayat-Ur-Rahman et al. (2007)



Table 4 continued

No.	Name	Plant origin	Family	Chem. formula	$M_{\rm r}$	References
22	Mauritine-K	Ziziphus mauritiana Lam.	Rhamnaceae	C <sub>31</sub> H <sub>47</sub> N <sub>5</sub> O <sub>5</sub>	569	Singh et al. (2007b)
23	Apetaline-A	Ziziphus apetala Hook. f.	Rhamnaceae	$C_{30}H_{35}N_5O_6$	561	Han et al. (2011)
24	Apetaline-B			$C_{32}H_{39}N_5O_5$	573	
25	Epimauritine-A			$C_{32}H_{41}N_5O_5$	575	
26	Epimauritine-A N-oxide			$C_{32}H_{41}N_5O_6$	591	
27	Mauritine-A N-oxide			$C_{32}H_{41}N_5O_6$	591	
28	Apetaline-C			$C_{32}H_{39}N_5O_6$	589	
29	Hymenocardine N-oxide	Hymenocardia acida Tul.	Phyllanthaceae	$C_{37}H_{50}N_6O_7$	690	Tuenter et al. (2016)
30	Hymenocardinol			$C_{37}H_{52}N_6O_6$	676	
31	Hymenocardine-H			$C_{34}H_{51}N_7O_6$	653	

### Neutral cyclopeptide alkaloids

Neutral cyclopeptide alkaloids follow the same structural pattern as the other cyclopeptide alkaloids, but the terminal basic amino acid known as building block A (Fig. 1) is replaced by a substituent that does not contain a nitrogen atom (or that is not derived from an amino acid), e.g. an acyl group. It should be noted that the absence of the nitrogen atom in this substituent is a less ambiguous criterion than the absence of basic properties, since N-formylation destroys its basic character. Nevertheless, *N*-formyl containing cyclopeptide alkaloids as described above should not be classified as "neutral cyclopeptide alkaloids" in order not to create confusion. Because the number of neutral cyclopeptide alkaloids is still rather small, there has been no particular need to distinguish subtypes, but obviously also here the same distinction between 13-, 14- and 15-membered ring types consisting of 4 or 5 building blocks (depending on the presence or absence of unit E) can be made. Two new neutral cyclopeptide alkaloids have been reported from the root of Ziziphus oxyphylla Edgew., i.e. oxyphylline-B (32) and oxyphylline-C (33) (Table 5), together with oxyphylline-D, nummularine-C and -R (Kaleem et al. 2012). Oxyphylline-B consists of proline, isoleucine and valine as building blocks B, C and E, respectively; oxyphylline-C of phenylalanine, proline and again phenylalanine, respectively. In both compounds the "neutral" or rather nitrogenlacking moiety is a cinnamoyl group. It should be noted that a compound obtained from Ziziphus oxyphylla that was named oxyphylline-D had already been reported before from *Sphaeranthus indicus* L. (Nisar et al. 2010). Also hemsine-A was reported from *Z. oxyphylla* (Choudhary et al. 2011).

In amaiouine (34), isolated from leaves of Amaioua guianensis Aubl. (Rubiaceae), units B, C and E are phenylalanine, phenylalanine and proline, respectively, in which the proline-N is again substituted with a cinnamoyl residue (Laurindo de Oliveira et al. 2009). Compounds 32–34 can therefore be considered as 5(14) neutral cyclopeptide alkaloids. Four new neutral cyclopeptide alkaloids of the 4(14) type have been isolated from root bark of Scutia buxifolia Reiss. (Rhamnaceae) (Maldaner et al. 2011), in addition to the known compounds scutianine-B, -C, -D and -E. Scutianene-E (35) and two of its diastereoisomers 3,4,28-tris-epi-scutianene-E (36) and 28-epi-scutianene-E (37) consist of  $\beta$ -hydroxy-phenylalanine as ring-bound amino acid (building block C), and leucine as building block B. A cinnamoyl moiety is directly attached to N-atom of the  $\beta$ -hydroxy-leucine unit. Similarly, in scutianene-L (38) isoleucine is the ringbound amino acid (unit C), whereas the β-hydroxyamino acid (unit B) is phenylalanine, the N-atom of which is substituted with a cinnamoyl group. A related 4(14) type compound, justicianene-A, was obtained from whole plant of Justicia procumbens L. (Acanthaceae), but interestingly a tyrosine moiety is present instead of a hydroxystyrylamine unit (Jin et al. 2015). This adds evidence to the hypothesis that the hydroxystyrylamine unit is biogenetically derived from tyrosine (Gournelis et al. 1997).

There may be some controversy whether or not these neutral cyclopeptides should be considered as



Table 5 Neutral cyclopeptides

$$R_2$$
 $NH-R_3$ 

$$\begin{tabular}{ll} \bf 32 \ Oxyphylline-B & R_1 & CH(CH_3)CH_2CH_3 \end{tabular}$$

 $R_2$   $CH(CH_3)_2$   $R_3$  Cinnamoyl

$$\begin{array}{c|c} & & & \\ & & \\ & & & \\ & & \\ & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

33 Oxyphylline-C  $R_1$  Benzyl

R<sub>2.</sub> Cinnamoyl

34 Amaouine

R<sub>1</sub> Benzyl

R<sub>2.</sub> Cinnamoyl

**35** Scutianene-E R<sub>1</sub> 1-OH-Benzyl (*L-threo*)

 $R_2$  CH(CH<sub>3</sub>)<sub>2</sub> (*L-erythro*)

R<sub>3</sub> Cinnamoyl

**36** 3,4,28-*tris-epi*-Scutianene R<sub>1</sub> 1-OH-Benzyl (*L-erythro*)

 $R_2$  CH(CH<sub>3</sub>)<sub>2</sub> (*D-erythro*)

R<sub>3</sub> Cinnamoyl

**37** 28-epi-Scutianene R<sub>1</sub> 1-OH-Benzyl (*L-erythro*)

 $R_2$  CH(CH<sub>3</sub>)<sub>2</sub> (L-erythro)

R<sub>3</sub> Cinnamoyl

 $\begin{array}{lll} \textbf{38} \; \text{Scutianene-L} & & \text{R}_1 & & \text{CH(CH}_3)\text{CH}_2\text{CH}_3 \\ \end{array}$ 

R<sub>2</sub> Benzyl (*D-threo*)

R<sub>3</sub> Cinnamoyl

$$R_2$$
 $NH$ 
 $O$ 
 $R_3$ 
 $R_1$ 

39 Justicianene-A R<sub>1</sub> Benzyl

 $R_2$   $CH(CH_3)_2$ 

R<sub>3</sub> Cinnamoyl

No.	Name	Plant origin	Family	Chem. formula	$M_{\rm r}$	References
32	Oxyphylline-B	Ziziphus oxyphylla Edgew.	Rhamnaceae	$C_{33}H_{40}N_4O_5$	572	Kaleem et al. (2012)
33	Oxyphylline-C			$C_{40}H_{38}N_4O_5$	654	
34	Amaiouine	Amaioua guianensis Aubl.	Rubiaceae	$C_{40}H_{38}N_4O_5$	654	Laurindo de Oliveira et al. (2009)
35	Scutianene-E	Scutia buxifolia Reiss.	Rhamnaceae	$C_{32}H_{33}N_3O_5$	539	Maldaner et al. (2011)
36	3,4,28- <i>tris-epi</i> -Scutianene-E			$C_{32}H_{33}N_3O_5$	539	



Table 5 continued

No.	Name	Plant origin	Family	Chem. formula	$M_{\rm r}$	References
37	28-epi-Scutianene-E			$C_{32}H_{33}N_3O_5$	539	
38	Scutianene-L			$C_{32}H_{32}N_3O_4$	523	
39	Justicianene-A	Justicia procumbens L.	Acanthaceae	$C_{33}H_{36}N_3O_6$	570	Jin et al. (2015)

alkaloids, since they are missing the peptidogenic amino acid with a mono- or dimethylated amino group exhibiting basic properties (unless in the case of formyl-substitution). However, many alkaloids devoid of a basic nitrogen are known, such as colchicine or piperine, and similarly also the neutral compounds discussed above could be termed "cyclopeptide alkaloids". Until 2011 only 8 neutral cyclopeptide alkaloids had been reported: Scutianene-C (=scutianene-D), discarenes-C and -D, discarines-M and -N, lotusanine-B, sanjoinenine and amaiouine (Maldaner et al. 2011). In the meantime oxyphyllines-B and -C, scutianene-E and its 2 isomers, scutianene-L and justitiacene-A have been added to this list, bringing the total to 15 representatives. All of them contain a 14-membered macrocyclic ring. Remarkably, all recently isolated neutral cyclopeptide alkaloids as listed in Table 5 contained a cinnamoyl moiety as the *N*-acyl group. It has been proposed by Maldaner et al. (2011) to use the ending *ine* for cyclopeptide alkaloids sensu stricto, and to use ene for neutral compounds.

## Distribution of cyclopeptide alkaloids

This review paper confirms that the Rhamnaceae family and especially the genus Ziziphus is by far the most important source of cyclopeptide alkaloids. Other plant families from which cyclopeptide alkaloids have been obtained during the past decade include the Acanthaceae, Malvaceae, Phyllanthaceae and Rubiaceae. The presence of the cyclopeptide alkaloids frangulanine and melonovine-A in Christiana africana DC. supported the reclassification of the genus Christiana from Tiliaceae to Malvaceae (Michalet et al. 2008). Recently UPLC/QTOF MS combined with an informatics platform was applied for the rapid characterisation of Ziziphi Spinosae Semen, the dried seeds of Ziziphus jujuba Mill. var. spinosa (Bunge) Hu ex H.F. Chou, and to distinguish it from its adulterant Ziziphi Mauritianae Semen (Zhang et al. 2016). With regard to cyclopeptide alkaloids, only ramosine-A, sanjoinine-A and lotusine-B were detected in Ziziphi Spinosae Semen, whereas a much wider range was present in Ziziphi Mauritianae Semen, including amphibine-D, lotusanine-A, lotusine-B, ramosine-A, sanjoinenine, sanjoinine-A, -B, -D, -F, G1 and its 11-epimer, and -G2. Daechuine S10 has been obtained from roots of *Z. jujuba* Mill. var. *spinosa* (Meng et al. 2013).

## Configurational studies and total synthesis

For some cyclopeptide alkaloids originally reported before 2006, the relative or absolute configuration has been investigated more recently. The relative configuration of oxyphylline-D, nummularin-C and -R was reported by Nisar et al. (2010). Single crystal X-ray diffraction of the 13-membered ring compound nummularine-B methiodide revealed all-S configurations of the amino acid residues (Panseeta et al. 2011). The absolute configuration of franganine, isolated from root bark of Discaria americana Gillies & Hook. (Rhamnaceae), was established by NMR spectroscopy and X-ray diffraction analysis of tri-N-methylfranganine methiodide (Caro et al. 2012). The absolute configuration of discarine-C, -D and myrianthine-A, three cyclopeptide alkaloids originally obtained from Discaria febrifuga, was determined by a combination of NMR studies, chiral gas chromatography, and comparison of NMR data with those of synthetic tripeptides (Mostardeiro et al. 2013).

Because of their diverse biological properties (see below), the cyclopeptide alkaloids and non-natural analogues have been of considerable interest to synthetic organic chemists. A two-step synthesis of p-cyclophanes (such as the 14-membered ring cyclopeptide alkaloids) by the combined use of a Ugi four-component reaction (Ugi-4CR) and an intramolecular  $S_N$ Ar-based macrocyclisation reaction has been reported as a sequence allowing the



introduction of at least 4 points of diversity (Cristau et al. 2006). De Greef et al. (2006) have developed a flexible two-step route to macrocyclic ansapeptoids and peptides, in which the core structure is synthesised by a combination of a Ugi four-component reaction with bifunctional building blocks to form the dipeptide part, followed by a suitable macrocyclisation reaction. The synthesis of cyclopeptide alkaloids starting from amino acids as building blocks and using copper(I) catalysis to install the key structural elements was discussed by Evano (2008). The total synthesis of the cyclopeptide alkaloids paliurine-E and -F, ziziphine-N and -Q, abyssenine-A and mucronine-E was reported, involving an intramolecular amidation of vinyl iodide, which allowed to address simultaneously two synthetic challenges associated with cyclopeptide alkaloids: The formation of the enamide and macrocyclisation (cycloenamidation reaction). Physical, spectroscopic and spectrometric characteristics of synthetic (-)-paliurine-F and mucronine-E corresponded in all respects to those reported for the natural products, thereby establishing their relative and absolute configurations (Toumi et al. 2007; 2008a, b; 2009). The total synthesis of abyssenine-B and mucronine-E was also reported by Wang et al. (2007) using a CuI/N,N-dimethylglycine-catalyzed coupling reaction of vinyl iodides with amides as the key step. The configuration of natural abyssenine-B and mucronine-E could tentatively be assigned as S,S,S. The total synthesis of ziziphine-N was also reported using a Mitsunobu reaction, followed by installation of the enamide part and ring closure (He et al. 2007). More recently Cu-mediated enamide formation in the total synthesis of complex peptide natural products was reviewed by Kuranaga et al. (2014).

### **Biological activity**

Activity on the Central Nervous System (CNS), analgesic and anti-inflammatory activity

As pointed out in previous reviews, many cyclopeptide alkaloids exert activity on the Central Nervous System (CNS) (Gournelis et al. 1997; El-Seedi et al. 2007). The cyclopeptide alkaloid fraction from Ziziphi Spinosi Semen, defined as the dried seed of *Ziziphus jujuba* Mill. var. *spinosa* (Rhamnaceae), and traditionally used

as a tranquiliser, analgesic and anticonvulsant, was found to enhance pentobarbital-induced sleeping behaviour in mice after oral administration. It was suggested that the enhancement of Cl<sup>-</sup> influx by the cyclopeptide alkaloid fraction may play an important role in the potentiation of pentobarbital-induced sleeping behaviour (Ma et al. 2008). The observed effects were comparable to those of muscimol used as a positive control. The cyclopeptide alkaloid fraction was prepared using a common liquid/liquid partition scheme for isolation of alkaloids, but unfortunately it was not phytochemically characterised. The same alkaloid fraction also showed anxiolytic effects: it increased the time spent on the open arms and the number of open arm entries in the elevated plus-maze test. Significant effects were obtained at a dose of 8.0 mg/kg, whereas diazepam used as a positive control was administered at a dose of 2.0 mg/kg. In addition the cyclopeptide alkaloid fraction increased the number of head-dips in the hole-board test (significant effect at 8.0 mg/kg), and increased the percentage of centre zone ambulatory time in the open-field box (significant effect at 2.0 mg/kg), again versus diazepam active at a dose of 0.5 and 2.0 mg/kg, respectively. However, in contrast to diazepam, it did not affect locomotor activity, and it did not influence grip force. The cyclopeptide alkaloid fraction was found to increase Cl influx and to overexpress y-subunits of GABAA receptors in cultured cerebellar granule cells (Han et al. 2008).

The cyclopeptide alkaloid fraction, prepared using a common liquid/liquid partition scheme for isolation of alkaloids from leaves of Ziziphus nummularia, was evaluated after oral administration for its analgesic activity in the acetic acid induced writhing, tail flick and hot plate tests; and for its anti-inflammatory activity against rat paw oedema, mouse peritonitis and cotton pellet granuloma. Although it was mentioned that the presence of cyclopeptide alkaloids was confirmed by identification tests, TLC and GC-MS analysis, there is no information on the composition of the extract. Anti-oedematogenic and anti-nociceptive effects were observed (Goyal et al. 2013). In the antioedematogenic assays significant effects were observed in a dose range of 10-30 mg/kg versus indomethacine used as a positive control at 10 mg/kg. Analgesic activity in the acetic acid-induced writhing test was observed in the same dose range versus aspirin as a positive control at 100 mg/kg. In addition, analgesic effects were also observed in the tail flick



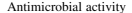
and hot plate tests, used to evaluate centrally acting analgesics, versus morphine at 5 mg/kg as a positive control. However, in an acute toxicity study, the  $LD_{50}$  value was established as 200 mg/kg, leaving a rather narrow therapeutic range.

The potential antinociceptive effect of six (4)14-membered cyclopeptide alkaloids, all belonging to the frangulanine-type, was investigated in mice using the tail-flick test as a simple pain model. Test compounds were administered intrathecally into the spinal column (Trevisan et al. 2009). Obviously results obtained after intrathecal administration cannot be compared to those observed after oral administration as in the studies mentioned above. Franganine and adouetine-X showed antinociceptive effects; adouetine-X also exhibited a pronounced analgesic effect in a chronic neuropathic pain model in mice, but unfortunately no positive control was used. Adouetine-X was able to decrease the activities of Ca<sup>2+</sup>-ATPase and Na<sup>+</sup>/K<sup>+</sup>-ATPase in vitro.

Five cyclopeptide alkaloids isolated from *Ziziphus oxyphylla*, including oxyphylline-B, -C, -D, nummularin-C and -R were evaluated in the acetic acid induced writhing and formalin induced flinching behaviour tests after intraperitoneal administration. Especially oxyphylline-B and nummularin-R showed activity. Significant activities were already observed at a dose of 2.5 mg/kg, versus diclofenac used as a positive control at 10 mg/kg. It was concluded that the peripheral analgesia was strongly augmented by their central effects (Kaleem et al. 2013b).

In summary, it appears that there is increasing evidence for the CNS activities of particular cyclopeptide alkaloids at realistic doses. In view of the limited number of cyclopeptide alkaloids evaluated for their effects on the CNS, it has not been possible yet to establish clear structure—activity relationships.

It has been reported that traditional practitioners of Indian medicine extract the stem part of *Ziziphus jujuba* by a crude pyrolysis method and use the oil in the treatment of pain. A prototype pyrolyser was applied to simulate this traditional method. FTIR and GC–MS analysis of the extracted oily substance obtained by both the traditional as the simulated process revealed the presence of various cyclic, nitrogenous, long chain and heterocyclic compounds, which were believed to be the pyrolysates of various cyclopeptide alkaloids present in the stem of *Ziziphus jujuba* (Shanmugavasan et al. 2011).



Mauritine-K, isolated from Ziziphus mauritiana Lam., exhibited antifungal activity (inhibition of spore germination) against some plant pathogenic fungi such as *Botrytis cinerea* at doses ranging from 200 to 1000 μg/ml, but since no positive control was used it is difficult to evaluate these results. Sativanine-K on the other hand was not active (Singh et al. 2007b). Bioassay-guided fractionation of a leaf extract of Melochia odorata L.f. (Malvaceae) resulted in the isolation of frangulanine, which showed moderate antifungal activity against Candida albicans and Saccharomyces cerevisiae. The minimal amount of frangulanine needed to inhibit fungal growth on a TLC plate was 25 and 50 µg, respectively, compared to 2.5 and 5.0 µg for ketoconazole, respectively (Emile et al. 2007).

Mauritine-L, -M, nummularine-H, -B and hemsine-A exhibited antiplasmodial activity against *Plasmodium falciparum* strain K1 with IC<sub>50</sub> values ranging from 3.7 to 10.3  $\mu$ M, whereas dihydroartemisinin used as a positive control exhibited an IC<sub>50</sub> value of 4.2 nM (Panseeta et al. 2011). Hymenocardine, its *N*-oxide, hymenocardine-H and hymenocardinol obtained from *Hymenocardia acida* showed antiplasmodial activity (strain K1) in a concentration range from 12.2 to 27.9  $\mu$ M, versus chloroquine used as a positive control with an IC<sub>50</sub> value of 0.2  $\mu$ M. However, only moderate cytotoxicities were observed against human lung fibroblasts (MRC-5 cells), yielding favourable selectivity indices (Tuenter et al. 2016).

Mauritine-M and nummularine-H showed antimycobacterial activity against Mycobacterium tuberculosis with MIC values of 72.8 and 4.5 μM, respectively, whereas the standard drugs isoniazid and kanamycin sulfate showed MIC values of 0.4 and 4.2 µM, respectively. No cytotoxicity controls were carried out, therefore the selectivity cannot be evaluated (Panseeta et al. 2011). The synthetic cyclopeptide alkaloids paliurine-E and -F, ziziphine-N and -Q, abyssenine-A and mucronine-E were evaluated for cytotoxicity against the human HT1080 tumoural cell line, and for antibacterial activity against a methicillin-resistant strain of Staphylococcus aureus, Bacillus anthracis and Escherichia coli. No significant antibacterial activity was observed, whereas paliurine-F, abyssenine-A and mucronine-E were moderately cytotoxic, displaying IC<sub>50</sub> values of 0.82, 1.03 and



0.68 mM, respectively. No positive controls were used (Toumi et al. 2009).

Again, as mentioned above for the CNS activity, the number of cyclopeptide alkaloids evaluated for their antimicrobial effects still is too low to allow the establishment of clear structure–activity relationships. Nevertheless, it appears that the cyclopeptide alkaloids as a class can be considered as a promising source of new antimicrobial lead compounds, especially against some fungi, mycobacteria or protozoa, with favourable selectivity indices, in the absence of obvious cytotoxicity.

## Other activities

Three cyclopeptide alkaloids isolated from Ziziphus oxyphylla Edgew., nummularine-C, -R and hemsine-A showed inhibition of  $\alpha$ -glucosidase and anti-glycation activities, which may support local antidiabetic use. With regard to α-glucosidase inhibition, all compounds were more active than 1-deoxynojirimycin, used as a positive control. Hemsine-A was more active as anti-glycation agent than rutine, used as a positive control in this assay (bovine serum albumin—methyl glyoxal assay) (Choudhary et al. 2011). Oxyphylline-D, nummularin-C and -R from the same species were found to be active as in vitro inhibitors of urease, catalysing the production of ammonia and carbon dioxide from urea, which plays a role in various pathologies. All test compounds were more active than thiourea used as a positive control (Kaleem et al. 2013a). Oxyphylline-B, -C, -D, nummularine-C and -R showed in vitro antioxidative (radical scavenging) potential (Kaleem et al. 2015).

For all biological activities, it should be noted that publications dealing with extracts, e.g. *Ziziphus* extracts, were only included in this review if at least the presence of cyclopeptide alkaloids had been confirmed. Reports dealing with biological activities of completely uncharacterised extracts were not included in this review.

# Conclusions

Although the number of cyclopeptide alkaloids is gradually increasing, it remains a relatively small class of natural products. All cyclopeptide alkaloids identified in the past decade follow the same structural

patterns as outlined before. Remarkably, from the 39 compounds reported for the first time from nature in this time frame, 8 belong to the class of neutral cyclopeptide alkaloids, bringing the total of representative of this group to 15. It is proposed to use the wording "neutral cyclopeptide alkaloid" only for those compounds containing a side that does not contain a nitrogen atom (or that is not derived from an amino acid), e.g. an acyl group. Indeed, the absence of the nitrogen atom in this substituent is a less ambiguous criterion than the absence of basic properties, since N-formylation of an amino-acid derived moiety results in loss of the basic character, but the latter compounds should not be considered as neutral cyclopeptide alkaloids. By analogy with the differentiation between 4(13), 5(13), 4(14) and 5(14) cyclopeptide alkaloids, it is proposed to distinguish two types of neutral cyclopeptide alkaloids, i.e. the 4(14) and the 5(14) type, in which the N-acyl moiety is considered as the fourth resp. the fifth building block.

When reviewing the literature it becomes obvious that on many occasions the relative or absolute configuration of all chiral centers has not been determined yet. This remains a challenge for future chemical and spectroscopic work. A second observation is that relatively few compounds have pharmacologically been investigated, and that even fewer compounds have been the subject of systematic investigations to establish structure—activity relationships. Small libraries of cyclopeptide alkaloids should be constructed (by isolation of synthetically) in order to fill this gap.

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#### References

Caro MSB, De Oliveira LH, Ilha V et al (2012) Absolute configuration of franganine. J Nat Prod 75:1220–1222

Choudhary MI, Adhikari A, Rasheed S et al (2011) Cyclopeptide alkaloids of *Ziziphus oxyphylla* Edgw as novel inhibitors of  $\alpha$ -glucosidase enzyme and protein glycation. Phytochem Lett 4:404–406

Cristau P, Vors JP, Zhu J (2006) Rapid synthesis of cyclopeptide alkaloid-like *para*-cyclophanes by combined use of Ugi-4CR and intramolecular S<sub>N</sub>Ar reaction. QSAR Comb Sci 25:519–526

De Greef M, Abeln S, Belkasmi K et al (2006) Rapid combinatorial access to macrocyclic ansapeptoids and



- ansapeptides with natural-product-like core structures. Synthesis 23:3997–4004
- Dias GCD, Gressler V, Hoenzel SCSM et al (2007) Constituents of the roots of *Melochia chamaedrys*. Phytochemistry 68:668–672
- El-Seedi HR, Larsson S, Backlund A (2005) Chemosystematic value of cyclopeptide alkaloids from *Heisteria nitida* (Olacaceae). Biochem Syst Ecol 33:831–839
- El-Seedi HR, Zahra MH, Goransson U et al (2007) Cyclopeptide alkaloids. Phytochem Rev 6:143–165
- Emile A, Waikedre J, Herrenknecht C et al (2007) Bioassyguided isolation of antifungal alkaloids from *Melochia odorata*. Phytoter Res 21:398–400
- Evano G (2008) Synthèse de produits naturels: des acides amines et du cuivre(I) pour la synthèse d'alcaloides cyclopeptidiques. Actualité Chimique 322:20–26
- Gournelis DC, Laskaris GG, Verpoorte R (1997) Cyclopeptide alkaloids. Nat Prod Rep 14:75–82
- Goyal M, Ghosh M, Nagori BP et al (2013) Analgesic and antiinflammatory studies of cyclopeptide alkaloid fraction of leaves of *Zizyphus nummularia*. Saudi J Biol Sci 20:365–371
- Han H, Ma Y, Eun JS et al (2008) Anxiolytic-like effects of cyclopeptide fraction alkaloids of Zizyphi Spinosi Semen: possible involvement of GABAA receptors. Biomol Ther 16:261–269
- Han J, Ji CJ, He WJ et al (2011) Cyclopeptide alkaloids from Ziziphus apetala. J Nat Prod 74:2571–2575
- He G, Wang J, Ma D (2007) Highly convergent route to cyclopeptide alkaloids. Total synthesis of ziziphine N. Org Lett 9:1367–1369
- Inayat-Ur-Rahman Khan MA, Arfan M et al (2007) A new 14-membered cyclopetide alkaloid from *Zizyphus oxy-phylla*. Nat Prod Res 21:243–253
- Jin H, Chen L, Tan Y et al (2015) New cyclopeptide alkaloid and lignin glycoside from *Justicia procumbens*. J Asian Nat Prod Res 17:33–39
- Joullié MM, Richard DJ (2004) Cyclopeptide alkaloids: chemistry and biology. Chem Commun 21:2011–2015
- Kaleem WA, Nisar M, Qayum M et al (2012) New 14-membered cyclopeptide alkaloids from Zizyphus oxyphylla Edgew. Int J Mol Sci 13:11520–11529
- Kaleem WA, Nisar M, Qayum M et al (2013a) Urease inhibitory potential of *Zizyphus oxyphylla* Edgew. extracts and isolated compounds. Turkish J Med Sci 43:497–500
- Kaleem WA, Muhammed N, Qayum M et al (2013b) Antinociceptive activity of cyclopeptide alkaloids isolated from Ziziphus oxyphylla Edgew (Rhamnaceae). Fitoterapia 91:154–158
- Kaleem WA, Muhammad N, Khan H et al (2015) Antioxidant potential of cyclopeptide alkaloids isolated from Zizyphus oxyphylla. J Chem Soc Pak 37:474–478
- Kang KB, Ming G, Kim GJ et al (2015) Jubanines F-J, cyclopeptide alkaloids from the roots of *Ziziphus jujuba*. Phytochemistry 119:90–95
- Kuranaga T, Sesoko Y, Inoue M (2014) Cu-mediated enamide formation in the total synthesis of complex peptide natural products. Nat Prod Rep 31:514–532
- Laurindo de Oliveira P, Tanaka CMA, Kato L et al (2009) Amaouine, a cyclopeptide alkaloid from the leaves of Amaioua guianensis. J Nat Prod 72:1195–1197

- Ma Y, Han H, Nam SY et al (2008) Cyclopeptide alkaloid fraction from Zizyphi Spinosi Semen enhances pentobarbital-induced sleeping behaviors. J Ethnopharmacol 117:318–324
- Maldaner G, Marangon P, Ilha V et al (2011) Cyclopeptide alkaloids from *Scutia buxifolia* Reiss. Phytochemistry 72:804–809
- Meng YJ, Zhang YW, Jiang HY et al (2013) Chemical constituents from the roots of *Zizyphus jujuba* Mill. Biochem Syst Ecol 50:182–186
- Michalet S, Payen-Fattaccioli L, Beney C et al (2008) New components including cyclopeptides from barks of *Christina Africana* DC. (Tiliaceae). Helv Chim Acta 91:1106–1117
- Morel AF, Maldaner G, Ilha V (2009) Cyclopeptide alkaloids from higher plants. In: Cordell G (ed) The alkaloids: chemical and biological perspectives, vol 67. Wiley, New York, p 79
- Mostardeiro MA, Ilha V, Dahmer J et al (2013) Cyclopeptide alkaloids: sterochemistry and synthesis of the precursors of discarines C and D and myrianthine A. J Nat Prod 76:1343–1350
- Nisar M, Kaleem WA, Adhikari A et al (2010) Sterochemistry and NMR data assignment of cyclopeptide alkaloids from *Zizyphus oxyphylla*. Nat Prod Commun 5:1205–1208
- Pandey MB, Singh AK, Singh JP et al (2008a) Three new cyclopeptide alkaloids from *Zizyphus* species. J Asian Nat Prod Res 10:709–713
- Pandey MB, Singh AK, Singh VP et al (2008b) Cyclopeptide alkaloids from *Zizyphus sativa* bark. Nat Prod Res 22:219–221
- Pandey MB, Singh JP, Singh AK et al (2008c) Xylopyrine-F, a new cyclopeptide alkaloid from *Zizyphus xylopyra*. J Asian Nat Prod Res 10:725–728
- Pandey MB, Singh JP, Singh S et al (2008d) Constituents of Zizyphus jujuba. J Indian Chem Soc 85:555–556
- Pandey MB, Singh S, Malhotra M et al (2012) Two new 14-membered cyclopeptide alkaloids from *Zizyphus xylopyra*. Nat Prod Res 26:836–841
- Panseeta P, Lomchoey K, Prabpai S et al (2011) Antiplasmodial and antimycobacterial cyclopeptide alkaloids from the root of Ziziphus mauritiana. Phytochemistry 72:909–915
- Rai N, Singh S, Singh VP et al (2008) Cyclopeptide alkaloids of Zizyphus xylopyra. J Indian Chem Soc 85:336–337
- Shanmugavasan S, Vaitheeswaran KSR, Ramachandran T (2011) Design and development of pyrolyser to extract medicinal oil from the stem of *Ziziphus jujuba*. J Anal Appl Pyrol 92:176–183
- Singh S, Pandey MB, Singh JP et al (2006) Peptide alkaloids from Zizyphus sativa bark. J Asian Nat Prod Res 8:733–737
- Singh AK, Pandey MB, Singh VP et al (2007a) Xylopyrine-A and xylopyrine-B, two new peptide alkaloids from *Zizy-phus xylopyra*. Nat Prod Res 21:1114–1120
- Singh AK, Pandey MB, Singh VP et al (2007b) Mauritine-K, a new antifungal cyclopeptide alkaloid from *Zizyphus mauritiana*. Nat Prod Res 21:781–784
- Singh AK, Pandey MB, Singh VP et al (2008a) Xylopyrine-C, a new cyclopeptide alkaloid from *Zizyphus xylopyra*. J Asian Nat Prod Res 10:715–718
- Singh A, Pandey MB, Singh JP et al (2008b) Peptide alkaloids of Zizyphus rugosa. J Indian Chem Soc 85:658–659



- Singh S, Pandey AK, Yadava A et al (2012a) Peptide alkaloids from *Zizyphus joazeiro*. J Chem Biol Phy Sci 2:608–611
- Singh S, Yadev A, Singh P (2012b) Chemical constituents of Zizyphus joazeiro. Int J Green Herbal Chem 1:18–20
- Singh JP, Raghubanshi S, Yadava A (2013) Cyclopeptide alkaloids of *Zizyphus rugosa*. Chem Biol Phys Sci 3:994–997
- Tan NH, Zhou J (2006) Plant cyclopeptides. Chem Rev 106:840–895
- Toumi M, Couty F, Evano G (2007) Total synthesis of paliurine F. Angew Chem Int Ed 46:572–575
- Toumi M, Couty F, Evano G (2008a) Total synthesis of the cyclopeptide alkaloid paliurine E. Insights into macrocyclisation by ene-enamide RCM. J Org Chem 72:9003–9009
- Toumi M, Couty F, Evano G (2008b) Eight-step total synthesis of the cyclopetide alkaloid mucronine E. Synlett 1:29–32
- Toumi M, Rincheval V, Young A et al (2009) A general route to cyclopeptide alkaloids: total synthesis and biological

- evaluation of paliurines E and F, ziziphines N and Q, abyssenine A, mucronine E, and analogues. Eur J Org Chem 20:3368–3386
- Trevisan G, Maldaner G, Velloso NA et al (2009) Antinociceptive effects of 14-membered cyclopeptide alkaloids. J Nat Prod 72:608–612
- Tuenter E, Exarchou V, Baldé A et al (2016) Antiplasmodial cyclopeptide alkaloids from root bark of *Hymenocardia acida*. J Nat Prod 79:1746–1751
- Wang J, Schaeffler L, He G et al (2007) Total synthesis and stereochemistry assignment of 15-membered peptide alkaloids abyssenine B and mucronine E. Tetrahedron Lett 48:6717–6721
- Zhang FX, Li M, Qiao LR et al (2016) Rapid characterization of Ziziphi Spinosae Semen by UPLC/Qtof MS with novel informatics platform and its application in evaluation of two seeds from Ziziphus species. J Pharm Biomed Anal 122:59–80

