Maculine: a furoquinoline alkaloid from the family Rutaceae: sources, syntheses and biological activities

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This paper is dedicated to Prof Jan Bergman for his 80th birthday and his involvement in organic chemistry, and with deep appreciation for over 40 years of friendship

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Abstract

Maculine is one of the furoquinolines which are characteristic of the family Rutaceae. Its chemical name is 9-methoxy[1,3]dioxolo[4,5-g]furo[2,3-b]quinoline and it was isolated from several genera of Rutaceae. The present mini-review covers the different sources of maculine, its methods of separation, synthetic pathways and biological activities.

Keywords: Maculine, furoquinoline, Rutaceae, alkaloids

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1. Introduction

The Rutaceae family has about 140 genera, consisting of herbs, shrubs and small trees which grow in all parts of the world, Moluccas, New Guinea, Australia, New Caledonia, Brazil, Cameroon. They are used in traditional medicine for treating snake bites, stomatitis, rheumatism, bronchitis and other diseases. Rutaceae plant is the source of furoquinoline alkaloids, besides furanocoumarin alkaloids, phenolic-structured compounds and terpenes. The main furoquinoline alkaloid compounds isolated from this family are dictamnine (1), skimmianine (2), γ-fagarine (3), haplopine (4), 7-hydroxydictamine (5), evolitnine (6), maculosidine (7), maculine (8), flindersiamine (9), tectaveardoornine (10), leptanoine A (11), leptanoine B (12), leptanoine C (13), melineurine (14), kokusaginine (15), acronydine (16), tecteamanienine A (17), tecteamanienine B (18), acronydine (19), isohaplopine (20), 5-(1,1-dimethylallyl)-8-hydroxyfuro[2,3-b]quinolone (21), isodictamnine (22), isomaculosidine (23), iso-γ-fagarine (24), dictangustine A (25), isopteleine (26), isoskimmianine (27), isokokusaginine (28), (+) isoplatydesmine (29), (+)-araliopsine (30), balfourodine (31), pseudobalfourodine (32), dihydroisodictammine (33), dihydridictamine (34), evoxine (35) and isoevoxine (36). Many of these compounds have antifungal, anti-bacterial, anti-Alzheimer, anti-plasmodial and phototoxic properties. Some of the others represent a promising perspective of being potential drugs. For example, it was reported that skimmianine (2), and melineurine (12) could potentially be used in the future to treat Alzheimer’s disease.
Figure 1. Structures of furoquinoline alkaloids isolated from Rutaceae.

The ability of γ-fagarine (3) to induce SCEs in human lymphocytes was demonstrated.\(^{23}\) Further, Nganou et. al. in 2019 reported the ability of maculine to be helpful in the fight against multi-drug resistant (MDR) cancer cells.\(^{24}\)

The present review highlights the sources, ways of extraction, and the methods of synthesis as well as the biological efficacy of maculine (8) as one of Rutaceae family alkaloids.
2.1. Botanical Sources

Maculine is a furoquinoline alkaloid widely spread in the bark of several Flindersia; a genus of 17 species of trees in the family Rutaceae. They grow naturally in the Moluccas, New Guinea, Australia, and New Caledonia.\textsuperscript{25-27} Also, maculine was isolated from the root bark of \textit{Araliopsis soyauxii} (family Rutaceae).\textsuperscript{13} Maculine was identified in the leaves and the fruits of \textit{Teclea nobilis}, a tropical African medicinal plant (Rutaceae).\textsuperscript{14} It was separated from the stem bark of \textit{Araliopsis tabouensis} Aubrev and Pellegr (Rutaceae); the large tropical west African tree whose bark used in the folk medicine for the treatment of gonorrhea.\textsuperscript{17} Maculine is abundant in \textit{Esenbeckia}, which is a genus of ca 30 species, native in tropical America.\textsuperscript{28} Species of this genus are known as a source of a variety of typical rutaceous secondary metabolites.\textsuperscript{29,30} Thus, maculine was isolated from \textit{E. almawillia} and \textit{E. grandiflora}\textsuperscript{9} and the leaves of \textit{Esenbeckia litoralis} collected from Temascal, Oaxaca, Mexico.\textsuperscript{31}

The phytochemical investigation of the hexane extract from the roots of \textit{Esenbeckia grandiflora} subsp. \textit{grandiflora var. grandiflora} (Rutaceae), native to tropical America allowed the identification of maculine.\textsuperscript{32} Furthermore, the methanol extract of the roots of \textit{Esenbeckia almawillia} (Rutaceae) was a source of maculine.\textsuperscript{33} \textit{Esenbeckia leiocarpa} (Rutaceae), popularly known as \textit{guarantã, goiabeira}, a native tree of Brazil, afforded maculine.\textsuperscript{16} Another plant of family Rutaceae, \textit{Oricia suaveolens} expressed maculine in the stems and leaves extract. This plant could be found in the Democratic Republic of the Congo, Ivory Coast, Ghana, Guinea, Nigeria, and Sierra Leone.\textsuperscript{34} The widespread nature of maculine in Rutaceae was further confirmed by investigation of \textit{Zanthoxylum buesgenii} and \textit{Teclea afzelii} which represented another source of this alkaloid.\textsuperscript{35} This last is a tropical African plant, distributed from Sierra Leone to Cameroon and used in Cameroonian folk medicine in the treatment of wound infections, abdominal pains, cough, fever and asthma.\textsuperscript{15} Recently maculine was isolated from the methanol extract of the stem bark of \textit{Araliopsis soyauxii} (Rutaceae).\textsuperscript{24}
Table 1. The various sources of masculine and its extraction from different parts of Rutaceae plants

<table>
<thead>
<tr>
<th>Genus</th>
<th>Species</th>
<th>Part of plant</th>
<th>Solvent of extraction</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Teclea</td>
<td>nobilis</td>
<td>Leaves and fruits</td>
<td>Ethanol</td>
<td>14</td>
</tr>
<tr>
<td>Teclea</td>
<td>nobilis</td>
<td>Roots</td>
<td>Dichloromethane: methanol (1:1)</td>
<td>35</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>methanol</td>
<td></td>
</tr>
<tr>
<td>Esenbeckia</td>
<td>leiocarpa</td>
<td>Stems</td>
<td>Ethanol</td>
<td>30</td>
</tr>
<tr>
<td>Esenbeckia</td>
<td>almawillia</td>
<td>Roots</td>
<td>Methanol</td>
<td>31</td>
</tr>
<tr>
<td>Teclea</td>
<td>afzelii</td>
<td>Stem bark</td>
<td>Methanol</td>
<td>15, 19</td>
</tr>
<tr>
<td>Flindersia</td>
<td>maculosa</td>
<td>Stem bark</td>
<td>Methanol</td>
<td>36, 37</td>
</tr>
<tr>
<td>Araliopsis</td>
<td>soyauxii</td>
<td>Root and stem bark</td>
<td>Petroleum ether and chloroform</td>
<td>19</td>
</tr>
<tr>
<td>Araliopsis</td>
<td>soyauxii</td>
<td>stem barks and leaves</td>
<td>Methanol</td>
<td>34</td>
</tr>
<tr>
<td>Zanthoxylum</td>
<td>buesgenii</td>
<td>Aerial parts</td>
<td>Dichloromethane: Methanol (1:1)</td>
<td>33</td>
</tr>
<tr>
<td>Araliopsis</td>
<td>tabouensis</td>
<td>Root and stem bark</td>
<td>Hexane and chloroform</td>
<td>17</td>
</tr>
<tr>
<td>Raulinoa</td>
<td>echinata</td>
<td>Stems and leaves</td>
<td>Hexane and MeOH</td>
<td>38</td>
</tr>
<tr>
<td>Esenbeckia</td>
<td>almawillia</td>
<td>Trunk bark</td>
<td>n-Hexane and EtOH</td>
<td>9</td>
</tr>
<tr>
<td>Esenbeckia</td>
<td>grandiflora</td>
<td>Roots</td>
<td>Ethanol, then n-hexane, CHCl₃ and EtOAc</td>
<td>9</td>
</tr>
<tr>
<td>Esenbeckia</td>
<td>litoralis</td>
<td>Leaves</td>
<td>Ethyl acetate</td>
<td>29</td>
</tr>
<tr>
<td>Vepris</td>
<td>punctata</td>
<td>Wood</td>
<td>CH₂Cl₂/MeOH</td>
<td>39</td>
</tr>
<tr>
<td>Oricia</td>
<td>suaveolens</td>
<td>Stems and leaves</td>
<td>Methanol</td>
<td>32</td>
</tr>
</tbody>
</table>

3. Organic Synthesis

Maculine has been synthesized in three different pathways. Firstly, it was prepared by condensation of 3,4-methylenedioxyaniline (37) with diethyl (2-benzylxoyethyl)malonate (38) to afford 3-(2-benzylxoyethyl)-4-hydroxy-6,7-methylenedioxy carbostyril (39). Compound 39, in turn, was methylated by diazomethane to provide the corresponding 4-methoxy compound 40. Cyclization of 40 by the action of polyphosphoric acid yielded 4-methoxy-6,7-methylenedioxy-2,3-dihydrofuro[2,3-b]quinoline (41) which produced the final product 8 (mp 195-196 °C) under dehydrogenation with N-bromosuccinimide (Scheme 1).
Scheme 1. Synthesis of Ohta and Mori.\textsuperscript{38} Reagents and conditions: (i) EtOH 2mol, diphenyl ether, heat, 30min, aq NaOH 5%, yield 45%; (ii) diazomethane, Et\(_2\)O, 0-5°C overnight, aq KOH 5%, yield 77%; (iii) polyphosphoric acid, heat, 1h, yield 31%; (iv) N-bromosuccinimide, AcOH, AcONa, CCl\(_4\), heat, 4h, aq NaOH 2%, yield 51%.

Zimmer and Walter\textsuperscript{43} reported a new pathway for synthesis of maculine via the utilization of certain substituted \(\alpha\)-benzylidene-\(\gamma\)-butyrolactones. Condensation of acetyl-butyrolactone (42) with 2-nitro-4,5-methylenedioxybenzoyl chloride (43) gave the corresponding 3-(6-nitro[1,3]benzodioxole-5-carbonyl)-dihydrofuran-2(3H)-one (44). The reaction of 44 with diazomethane produced (\(E\))-3-[methoxy-3-(6-nitro[1,3]benzodioxid-5-yl)methylene]dihydrofuran-2(3H)-one (45). Upon reduction of NO\(_2\) group of 45 using Pd/C in presence of methanolic HCl led to the formation of dihydromaculine 41. Aromatization of 41 upon refluxing and irradiating using UV-light in presence of an equivalent amount of \(N\)-bromosuccinimide in CCl\(_4\), followed by refluxing the residue with collidine gave 8\textsuperscript{43} (Scheme 2). Kuwayama\textsuperscript{44} has used a similar route using magnesium, CCl\(_4\), ethanol instead of sodium in the first step, and sodium methoxide for the aromatization of 41 instead of collidine.

Scheme 2. Synthesis of Zimmer and Walter.\textsuperscript{43} Reagents and conditions: (i) Na, MeOH yield 36%; (ii) diazomethane, C\(_6\)H\(_6\), 0-5°C, 5 days yield 91%; (iii) Pd/C, methanolic-HCl yield 72%; (iv) a: NBS, CCl\(_4\), reflux, 30-40min, UV-light, b: collidine, reflux.
In the last reported pathway, Yazima and Munakata demonstrated the synthesis of maculine starting with ethyl 2-[(1,3)benzodioxol-5-ylamino)-4-oxo-4,5-dihydrofuran-3-carboxylate (47). Thermal ring closure of the ester 47 under fusion led to the formation of [1,3]dioxolo[4,5-g]furo[2,3-b]quinoline-8,9(5H,7H)-dione (48). Methylation of 48 by reaction with diazomethane in dioxane yielded 49. Reduction of the furanone 49 using sodium borohydride gave the corresponding furanol 50. Dehydration of 50 furnished masculine (8) in 40% yield. 45 (Scheme 3)

Scheme 3: Synthesis of Yazima and Munakata. Reagents and conditions: (i) a: compound 37, TEA, r.t., dry THF, stirring, 40min, b: compound 46, Na, dry DHF, 0°C, α-chloropropionic acid, stirring, 40min, c: mixture of a and b with a, reflux, 20min, yield 10%; (ii) heat at 250°C, vigorous stirring, 20min, inert gas, yield 91%; (iii) diazomethane, dioxane, yield 40%; (iv) dry methanol, dry THF, sodium borohydride, stirring, 0°C, yield 31%; (v) dry dioxane, fused potassium bisulfate, refluxed, 45min, yield 40%.

4. Biological Activity

Although maculine showed varieties of biological activities that ranged from antimicrobial to antitumor, it has not yet been reported to possess a significant activity toward some particular targets. It revealed a phototoxic activity to certain yeasts (Saccharomyces cerevisiae), fungi (Candida albicans), and bacteria (Streptococcus faecalis) in long wave UV light. 20 This antimicrobial activity was further investigated by the disk diffusion method against Staphylococcus aureus in which the inhibition zone was moderate (9 mm) compared to the standard Ceftriaxone (20 mm). 36 Certain strains of bacteria and fungi were subjected to additional antimicrobial assays of maculine. The recorded antimicrobial activity in this assay was 87.5% against Escherichia coli, Pseudomonas aeruginosa, Staphylococcus aureus, Bacillus subtilis, Candida albicans, Candida gabrata, Salmonella typhimurum and Microsporum audorium. 46

The maximum efficacy as an antibacterial agent was observed against Bacillus subtilis. It recorded minimal microbial concentration MMC of 2.44 µg/ml (two-fold) lower than the reference gentamicin (MMC 4.22 µg/ml). Besides, it showed moderate activity against mycobacteria M. smegmatis with a minimal inhibition concentration MIC of 156.25 µg/ml compared to Ciprofloxacin. 46 The antiparasitic characteristic of maculine was studied against trypomastigote forms of Trypanosoma cruzi and malaria and it was moderate for both parasites. 19,47
Additionally, this furoquinoline alkaloid revealed a photo-mutagenic effect against a mutant strain of *Chlamydomonas reinhardtii* using a dose of 40 µg/ml with exposure of the cells to irradiation with UV-A.\(^{10,48}\) Its photosensitizer activity could be attributed to the planar structure, and so its capability of intercalating into DNA.\(^{48}\) Maculine displayed selective and moderate cytotoxic activity, and the data highlighted the possibility of using maculine to fight drug-sensitive and resistant cancers.\(^{24}\) It was tested on A2780 human ovarian cancer cell and the cytotoxicity was moderate of IC\(_{50}\) value = 4.2 µg/ml.\(^{40}\) Recently maculine was screened *in vitro* for its effect on the viability of two different human cancer cell lines, namely prostate PC-3 adenocarcinoma cells and colorectal HT-29 adenocarcinoma cells. The results revealed that maculine showed some antiproliferative effect, but exclusively on HT-29 cells at 100 µM.\(^{42}\)

### 5. Conclusions

Maculine is a furoquinoline alkaloid which is abundant in the Rutaceae family. Many species subclassified under this family have introduced a resource of this alkaloid. It could be noticed the lack of interest in the searching of analogues of maculine or even to develop more alternative synthetic procedures for its fabrication. We tried in this mini-review to draw attention toward this nitrogenous compound which however has not shown a breakthrough regarding the biological activity it could act as an aspect for drug discovery *via* structure optimization or as an inspiring scaffold.

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