The history of extensive structural modifications of pyrethroids

Kazuya Ujihara*

*Present address: Preferred Networks, Inc.,
Otemachi Bldg. 1–6–1 Otemachi, Chiyoda-ku, Tokyo 100–0004, Japan

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To whom correspondence should be addressed.
E-mail: ujihara@preferred.jp
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Fig. 1. Structure of pyrethrin I and the structure proposed by Stäudinger.

Introduction

Tanacetum cinerariifolium, commonly called an “insecticidal flower,” is one of three pyrethrum species and has been used as insect powder in the Dalmatian region since the Middle Ages due to its insecticidal activity in the ovule. Active ingredients extracted from dried pyrethrum, named pyrethrins, were among the few insect control agents available in the world before the invention of synthetic insecticides, but they were inconvenient for practical use.1) However, with the invention of a mosquito stick in 1890 (a mosquito coil in 1895) by Eiichiro Ueyama, a founder of the Japanese company Dainippon Jochugiku (Kincho), the use of pyrethrins gradually spread throughout the world due to its advantages in controlling flying insects and its ease of use. The mosquito coil is still one of the most popular materials for preventing mosquito bites without changing the coil shape. Even now, there is a demand for pyrethrins as natural insecticides, and 10,000 tons of dried petals are produced annually.2)

Six pyrethrins have been identified as insecticidal compounds in nature, and pyrethrin I (Fig. 1) is the most important component because of its overall insecticidal efficacy and abundance. Pyrethrin I has a unique ester structure constructed from substituted cyclopentenolone (called pyrethrolone) and a substituted cyclopropanecarboxylic acid (called chrysanthemic acid). Pyrethrins have fast action (i.e., a fast-acting effect), called a knockdown effect, which paralyzes insect pests by modifying the kinetics of the voltage-sensitive sodium channel.3) Additionally, they are active ingredients in mosquito coils, which are safe to mammals and have excellent heat transpiration. However, since pyrethrin I is constructed from many unstable partial structures, such as a trialkyl-substituted double bond, cyclopentenolone, and conjugated diene, the use of pyrethrins as agricultural insecticides has been very limited. Furthermore, since the supply of natural pyrethrins was sometimes unreliable, the industrial production of pyrethrins was expected but proved difficult because of structural complications. In order to solve these problems, extensive structural modifications of pyrethrins have been conducted for over half a century, and a number of derivatives, the so-called synthetic pyrethroids, have been born.

Figure 2 shows the brief history of the structural modifications of pyrethroids. The structural modifications of both alcohol and acid moieties in natural pyrethrins have led to a number of synthetic pyrethroids with diverse structural features. Many researchers have participated in the competition to develop superior new pyrethroids. As a result, a variety of structural modifications have been achieved, and more than thirty compounds are on the market. In this review, the historical flow of the struc-
1. Early Studies of Natural Pyrethrin Structures

As Japan had a big share in the pyrethrum market in the early 1900s, many of the early pyrethrum studies were conducted in Japan. The first isolation of pyrethrins was reported by Fujitani. He claimed in the report that pyrethrins are ester compounds. In 1923, Yamamoto reported that the acid part of pyrethrins is 3-butenyl-2,2-dimethylcyclopropanecarboxylic acid. The following year, two prestigious chemists, Stäudinger and Ružička at Eidgenössische Technische Hochschule Zürich (ETH), published a paper of several hundred pages reporting detailed structural studies. The pyrethrin I structure they proposed is shown in Fig. 1. Their works were great achievements in natural product chemistry in light of the scientific level at that time (there were no spectroscopic analytical instruments), although we now know that the proposed structure contained some mistakes. The correct planar structure was elucidated in 1945, and a complete structure, including stereochemistry, was determined in 1958; this became the basis for subsequent developments of pyrethroid chemistry. The first synthetic pyrethroid, allethrin, was invented at the United States Department of Agriculture (USDA) in 1949, and its patent was opened to the public the following year. Several companies, including Sumitomo Chemical, successfully commercialized allethrin and got into the pyrethroid business in 1953. It is notable that a complete stereochemistry of pyrethrin I was determined five years after the launching of allethrin. This means that the researchers did not know which isomer was active when allethrin was launched.

2. Structural Modification of the Alcohol Part of Pyrethrin I

Most chemists wanted to convert the complicated pyrethrolone, the alcohol part of pyrethrin I, to a simpler one. Several attempts are shown in Fig. 3. The first successful modification achieved was allethrin synthesis, in which one vinyl group was removed from the unstable diene structure. Allethrin exhibited knockdown activity comparable to that of pyrethrin I (i.e., fast action). Even now allethrin is widely used as an active ingredient in mosquito killers such as mosquito coils and aerosol spray. Replacement of the allyl group of allethrin with a propargyl group was previously evaluated by the USDA in 1961, but the efficacy was only 60% that of allethrin. In contrast, Dainippon Jochugiku reported that the propargyl derivative was 1.2 times more potent than allethrin. They proposed that the difference might be due to the difference in sample purity and/or the different bioassay methods. Subsequently, Sumitomo Chemical established a chiral synthesis of this moiety and succeeded in commercializing it as prallethrin. Prallethrin was the best cyclopentenolone-type pyrethroid, especially in terms of the

† Four digits enclosed in parentheses indicate the year of the priority date of the patent or the publication date of the journal in which each compound was originally described.
knockdown effect on various house pests, such as mosquitoes, flies, and cockroaches.

The synthetic variations of chrysanthemate have already been reported by Staudinger and Ružička in a paper on the structural elucidation of natural pyrethrins. They found that substituted benzyl ester and some unsaturated aliphatic alcohol esters have insecticidal activity. Barthel et al. of the USDA and McLaughlin Gormley King Co. (MGK) randomly screened various benzyl esters to discover barthrin and dimethrin (Fig. 3). These compounds have a potent killing effect against houseflies and are more easily synthesized than pyrethrin I, but their knockdown activities against houseflies were only one-tenth that of pyrethrin I’s. This work encouraged many researchers to perform further modifications. The first breakthrough was realized as a carbodiimide-N-hydroxymethylol ester, tetramethrin, via a phthalimide-N-hydroxymethylol ester. Tetramethrin showed faster knockdown action against houseflies than allethrin and pyrethrins, with lower oral acute toxicity on mice. This finding led to the invention of imiprothrin, which was registered in 1996 as a very fast knockdown agent against crawling insects, especially cockroaches.

The paragraph above focused on the fast action of pyrethroids. Although all of these compounds do not necessarily have an excellent killing effect on insects, barthrin and dimethrin have some killing effect, but it is inferior to that of pyrethrins. These studies were followed by that of Elliott et al. of the National Research Development Corp. (NRDC) (Fig. 4). As a result, a 5-benzyl-3-furfuryl alcohol ester, resmethrin, was invented via a para-allyl- or para-benzyl-benzyl alcohol ester. However, it is very difficult to synthesize as compared with the substituted benzyl esters invented at that time. As resmethrin has an outstanding lethal effect against various insects, it was commercialized as a household insecticide and is still one of the most important mosquito-killing agents. Independently,
Dainippon Jochugiku found that 2-furfuryl alcohol esters also have significant insecticidal activity. In particular, furame-thrin, in which the allyl group was replaced by the propargyl group, exhibited superior performance as an active ingredient in an electric vaporizer. Another important development was the Sumitomo Chemical discovery of phenothrin, in which the furan ring of resmethrin was replaced by a benzene ring. This replacement made synthesis easier and improved photostability. Because of these properties, 3-phenoxybenzyl alcohol would be used in many pyrethroids that were invented subsequently. The NRDC had already evaluated para-benzyl-substituted benzyl alcohol ester in 1965. Since they did not evaluate meta-substituted ones at that time, they lost the patent right. Why did they lag behind Sumitomo in the discovery of phenothrin? They already had data indicating that meta-allylbenzyl and para-phenoxybenzyl esters exhibit only weak activity compared to para-allyl and para-benzylbenzyl esters, respectively. Accordingly, they might have concluded that para-substitution and the methylene bridge between the benzene and benzy1 groups are very important in eliciting insecticidal activity.

Around the same time, BASF introduced an ethynyl group at the α-position of various benzyl-type pyrethroids such as dime-thrin and found enhanced potency as compared to that of unsubstituted derivatives (Fig. 5). This finding was immediately applied to various known pyrethroids, as in the case of S-2684 derived from phenothrin. However, this conversion had not been applied in the case of resmethrin and was highly limited in the case of phenothrin. On the other hand, replacing the ethynyl group in S-2684 with a cyano group improved insecticidal efficacy. The derivative of S2684 was commercialized as cyphenothrin, which was mainly used as an active ingredient in cockroach killers. Cyphenothrin is also recognized as the first Type II pyrethroid, which is generally characterized by the presence of an α-cyano group, as compared to the former pyrethroids, called Type I pyrethroids. Most pyrethroids that have been developed for agricultural use to date contain a cyano group, as described later. Type II pyrethroids not only improve killing efficacy but also have a different effect on sodium channels. Type I pyrethroids tend to activate sodium channels, while Type II pyrethroids prolong the activated state of channels.

![Fig. 5. Introduction of an ethynyl/cyano group into the α-position of alcohol.](image)

![Fig. 6. Structural modifications of cyphenothrin for use as a crop protection agent.](image)
3. Challenge to Develop an Active Ingredient for Crop Protection

Cyphenothrin’s lethal action is superior to that of the former pyrethroids, although the agricultural use of this reagent has been very limited. What are the drawbacks of cyphenothrin for agricultural use? First, the manufacturing cost of the acid part, especially construction of the cyclopropane ring, is too expensive as compared with the improvement in crop yield. Second, trisubstituted double bonds in the acid part are not stable against oxygen and sunlight. Third, the insecticidal spectrum is not good against insects living in the field. It was expected that these problems might be solved by replacing the acid part with some inexpensive and stable substructures.

Figure 6 shows three successful approaches to overcoming the issues above. The first invented compound was fenpropathrin, the design of which was based on the finding at the University of Tokyo that the tetramethylcyclopropanecarboxylic acid used in tetralethrin could be used as a pyrethroid acid to increase the stability against oxygen and sunlight. Fenpropathrin was effective against several agricultural pests, especially mite pests in fruit. Fenpropathrin was later launched in a niche market.

The second approach is to replace the cyclopropane ring with something inexpensive to reduce the synthetic cost. This was accomplished by opening the cyclopropane ring and replacing the remaining part with a substituted benzene ring, resulting in fenvalerate. Fenpropathrin was later launched in a niche market.

The third approach is to substitute the methyl groups connecting the double bond to halogen atoms to reduce photooxidative instability. This acid part was first reported by Farkaš et al. in 1958. They synthesized the allethrin analog and achieved the enhancement of insecticidal activity. However, their research was not well known and was discontinued because the photostable alcohol part having high lethality activity was not known when they did their research. However, a combination of this acid with the alcohol part of fenothrin and cyphenothrin made permethrin, cypermethrin, and deltamethrin, which exhibited a new order of insecticidal activity with photostability. These analogues were highly useful, not only in the field of household insecticides but also as agricultural insecticides. The remaining task in the industry was to find economical methods of synthesizing this unique acid, permethric acid. To that end, many industrial laboratories have joined the race to look for a cheap process of synthesizing permethric acid.

The method of synthesis described in Elliott’s patent is summarized in Fig. 7. This process is superior on a laboratory level, but there are several problems to be overcome: i) several steps are required to manufacture the starting material, which is unfavorable to making cheap agricultural insecticides; ii) it contains a risky reaction, such as ozone oxidation; and iii) it contains low atom economy reactions, such as that of carbon tetrachloride with triphenylphosphine. Among the various synthetic methods that were attempted in an effort to overcome these issues in many companies and institutes, the process (shown in Fig. 8) used by the Sagami Chemical Research Center was the winner of this synthetic race. The condensation of prenyl alcohol and orthoacetate ester caused the Claisen rearrangement to make 3,3-dimethyl-4-pentenoate. The obtained product was condensed with carbon tetrachloride and cyclized by treating with a base to give a molecular target. This process is one of the superior industrial synthesis processes because starting materials are cheap except for prenol and the process is constructed by short steps, redox-neutral reaction, and high atom economy (byproducts: methanol and the salt of hydrogen chloride). Later, the problem of industrializing prenol synthesis was successfully solved by Kuraray.

4. Fluorine Chemistry in Structural Modifications of Pyrethroids

The application of fluorine chemistry in studies for the discovery of pharmaceuticals and agrochemicals had become popular in the 1970s. Fluorine chemistry also successfully applied the structural modifications of pyrethroid chemistry (Fig. 9). The first example of success was cyfluthrin. Bayer revealed that only the 4-position of the benzyl group in cypermethrin need-
Another remarkable application of fluorine chemistry to pyrethroids involved polyfluorobenzyl esters (Fig. 10). The first polyfluorobenzyl ester pyrethroid may have been reported by Elliott. He described the synthesis of pentafluorobenzyl chrysanthemate in his patent document. Although the results of insecticidal tests of several alcohol esters were described in the patent, no result for a pentafluorobenzyl ester was described in it. It is likely that his research focused on the 4-allylbenzyl alcohol ester and that the ester was only used as a standard in the gas chromatography analysis. Sumitomo Chemical reported that pentachlorobenzyl tetramethylcyclopropanecarboxylate exhibits activity comparable to that of existing pyrethroids, but their studies appeared to be focused on the tetramethylcyclopropanecarboxylic acid part. Ten years later, Bayer found that permethric acid pentafluorobenzyl ester exhibits superior activity against various insects. trans-Isomers such as fenfluthrin exhibited especially fast action against dipterous insects. On the other hand, cis-isomers, such as NAK1901, were useful for controlling soil insects due to their higher volatility, higher hydrophilicity, and higher stability in soil bacteria relative to other pyrethroids.
Fenfluthrin and NAK1901 were abandoned in the late 1970s for economic reasons and/or due to their high susceptibility to nucleophilic attack at the 4-fluorine in the benzyl group. However, ICI invented tefluthrin by replacing the 4-fluorine with a methyl group as a soil insect control agent in combination with the acid part of cyhalothrin. On the other hand, Bayer replaced the 4-fluorine in fenfluthrin with a hydrogen atom to develop transfluthrin as a household insecticide.

The author's group removed one methyl group from the side chain, called norchrysantemic acid, as a smaller molecular structure while maintaining insecticidal activity. A 4-methoxymethyl derivative, metofluthrin, exhibited extremely high knockdown activity in vapor action against various insects, especially mosquitoes. A 4-methyl derivative, profluthrin, had an insecticidal effect against various fabric insects that was superior to that of traditional moth proofers. Momfluorothrin was invented by the introduction of a cyano group into the acid part of metofluthrin, which exhibited excellent knockdown efficacy against not only houseflies but also German cockroaches. One potential reason for these activities may be that the cyano group forms a hydrogen bond to the channel protein, facilitating noncovalent interactions with the biological target.

5. Beyond Cyclopropane and Ester Frameworks

The invention of fenvalerate made a breakthrough in pyrethroid chemistry by escaping 2,2-dimethylcyclopropanecarboxylic acid ester dogma, as mentioned above. Further, drastic structural modifications were inspired by this finding (Fig. 11). The first example was inspired by the similarity of the acid part of fenvalerate to valine. Zoecon replaced the acid part of fenvalerate with N-aryl valine, which is now known as fluvinate. Fluvinate has very low toxicity to honeybees. Another attempt was made to form a hybrid of DDT and pyrethroid structures. The Commonwealth Scientific and Industrial Research Organisation (CSIRO) had originally discovered DDT analogues such as GH74 and CP51543 during a structure–activity relationship study of DDTs intended to overcome their environmental problem. However, these compounds were abandoned because of the 1974 energy crisis. Cycloprothrin that is a hybrid compound CP51543 (acid part) and pyrethroid (alcohol part) exhibited high insecticidal activity with low fish and daphnia toxicity.

The low aquatic organism toxicity is an important property of insecticides for paddy rice fields. In this sense, cycloprothrin was the first pyrethroid suitable for use in paddy rice fields.

While drastic structural modifications had been attempted to overcome the ester framework, the derivatives elicited only weak activities. Shell reported that ether compounds inspired from the fenvalerate structure exhibit some insecticidal activities. However, their study was withdrawn because the potencies of their compounds were nearly 1/100 that of fenvalerate. The first breakthrough from this approach was SD47443, in which oxime ether was considered as a replacement for the ester framework. SD47443 exhibited insecticidal activity comparable to that of typical pyrethroids. Meanwhile, Nissan Chemical found β-gem-dimethyl-β-phenylpropionic acid to be applicable as a pyrethroid acid, even though similar structures such as α-gem-dimethylphenylacetates and α-gem-dimethyl-β-phenylpropionates were nearly inactive. Mitsui Toatsu found that gem-dimethyl derivatives lacking the carbonyl group of the ester linkage (namely, converting the ester linkage to the ether linkage) exhibited excellent insecticidal activity. This finding led to the development of etofenprox, which had not only broad-spectrum insecticidal activity but also low fish toxicity. Silafluofen, invented by Dainippon Jochugiku, was the first commercialized organosilicon insecticide that was driven by changing three substructures of etofenprox: i) replacing quarterly carbon with a silicon atom, ii) replacing ether oxygen with a methylene group, and iii) introducing a fluorine atom at the 4-position of the benzyl group that was inspired from the cyflu-thrin structure. Substitution with a silicon atom slightly reduced

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Fig. 11. Development of non-dimethylcyclopropanecarboxylic acid esters.
its insecticidal potency but was very effective at reducing fish toxicity.\textsuperscript{80} Silafluofen is the only pyrethroid compound with a rank A classification in the Japanese fish toxicity classification system.

6. Application of Pyrethroids for Specialized Uses

The structural diversity of pyrethroids is very great, and each compound exhibits characteristic biological properties, some of which have been commercialized for specific applications (Fig. 12). Empenthrin, which is currently the most volatile commercialized pyrethroid,\textsuperscript{81} has been widely used as an active ingredient in moth proofers.\textsuperscript{82} Empenthrin was inspired by the $\alpha$-ethynyl moiety of furamethrin obtained in the course of studies on the synthesis of furamethrin, even though $\alpha$-ethynyl-benzyl esters had already been reported by BASF, as mentioned above.\textsuperscript{83} Flumethrin is structurally characterized as having the (E)-chlorine in cyfluthrin replaced by a 4-chlorophenyl group.\textsuperscript{84} Since flumethrin exhibits specific activity against parasitic insects and ticks on various cattle, it is used as active ingredient in veterinary medicine.\textsuperscript{85} Acrinathrin, in which structurally unique (Z)-acrylic acid exists in the acid side chain, exhibits high miticidal activity.\textsuperscript{86} Pyrethroids have a broad spectrum of insecticidal activity and sometimes tend to cause resurgence, but this is less likely with acrinathrin because the overall insecticidal activity of acrinathrin is lower than that of typical pyrethroids.\textsuperscript{87}

Conclusion

A synthetic pyrethroid, allethrin, was invented by the USDA and commercialized by Sumitomo Chemical in 1953. Since then, a large number of researchers have joined the race to discover superior pyrethroid insecticides. Each structural transformation described in this article is a historically inevitable flow of ideas rather than the inventions of geniuses. Pyrethroids developed mainly for agricultural use are shown in Fig. 13 in order of their year of invention. As seen in this figure, many pyrethroids for agricultural use have entered the market since the commercialization of fenvalerate. All of them were invented between 1971 and 1984. This means that even great and structurally convertible compounds such as pyrethrin I were effective as lead comp-

![Fig. 12. Pyrethroids containing unique structures.](image)

![Fig. 13. Leading pyrethroids ordered by year of invention (for agricultural use after cyphenothrin).](image)
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