Lipid Excipients for Cannabinoid Drug Products
ABOUT GATTEFOSSÉ

Gattefossé provides innovative excipients and drug delivery solutions to health industries worldwide. Simplifying formulation decisions and shortening the drug development path are at the heart of the Gattefossé product offer.

Our product list includes solubilizers; penetration, permeation, and bioavailability enhancers; modified and sustained-release matrix formers; and inert vehicles for oral, topical, injectable, and mucosal routes of administration.

Each excipient is fully characterized by physiochemical properties and safety profile. In addition we provide guidance documents for formulation design in preclinical as well as late development stages.

Gattefossé ensures timely technical support to its customers with highly trained technical staff and a well-established network of affiliates and distributors in 70 countries.
This brochure provides an overview of the formulation opportunities with Gattechossé excipients in developing medicinal cannabinoids for oral, topical, transdermal, and alternative routes of administration.

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The introduction of the cannabis plant into Western medicine dates to the 19th century when the plant was first used as an additive to patented medicines. The development of cannabis based therapies in the modern sense however came to a halt in the 1970’s after the passing of laws that have since categorized cannabis as a scheduled (narcotic) substance.

Amidst the regulatory constraints, over one hundred active molecules (phytocannabinoids) of the plant have been identified. Synthetic forms of cannabinoids such as dronabinol, nabilone, and cannabidiol have been developed and marketed for treatment of nausea associated with chemotherapy and for myoclonic seizures in patients two years of age and older.

However, it was not until 2018 that the US FDA approved Epidiolex®, the first purified phytocannabinoid isolated from the cannabis plant. Epidiolex® is indicated for severe forms of epilepsy, namely Lennox-Gastaut and Dravet syndromes (Table 1).

Table 1. Marketed cannabinoid drugs as of 2018

<table>
<thead>
<tr>
<th>Marketed Drug</th>
<th>Indication</th>
<th>Dosage Form</th>
<th>API Molecule</th>
<th>Drug Strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epidiolex®</td>
<td>Myoclonic seizure</td>
<td>Oral Solution</td>
<td>Cannabidiol</td>
<td>100 mg/mL</td>
</tr>
<tr>
<td>Marinol®</td>
<td>Nausea and emesis associated with chemotherapy</td>
<td>Soft Gel Capsules</td>
<td>Dronabinol</td>
<td>10 mg</td>
</tr>
<tr>
<td>Syndros®</td>
<td>Nausea and vomiting associated with cancer</td>
<td>Oral Solution</td>
<td>Dronabinol</td>
<td>5 mg/mL</td>
</tr>
<tr>
<td>Cesamet™</td>
<td>Nausea and vomiting associated with cancer</td>
<td>Oral Capsule</td>
<td>Nabilone</td>
<td>1 mg</td>
</tr>
<tr>
<td>Sativex®</td>
<td>Neuropathic pain from Multiple Sclerosis (MS) and for intractable cancer pain</td>
<td>Buccal Spray</td>
<td>Cannabinoid; Dronabinol</td>
<td>25 and 27 mg/mL</td>
</tr>
</tbody>
</table>

With an enhanced understanding of the medicinal benefits and looming ease of regulations around the globe, there is an expressed need for science based, adequately dosed, and age appropriate cannabinoid medicines.

The benefits of medicinal cannabinoids include:

- Controlled and measured dosing
- Age appropriate, patient-centric dosage forms
- Alternatives to unhealthy modes of intake such as vaping or smoking
Among the hundreds of phytocannabinoids and terpenes isolated from cannabis, two very lipophilic molecules stand out: tetrahydrocannabinol (THC) and cannabidiol (CBD). Amounting between 30% and 50% of a single plant’s weight, they are known to affect physiological processes such as mood, memory, appetite, and pain sensation. They function by activating the endocannabinoid receptors of the nervous system which are found throughout the body, including in the brain.

THC and CBD have been extensively studied and are found to have unique qualities of their own.

THC is responsible for the euphoric and psychoactive effects of cannabis and has applications as medicine for treatment of chronic pain and nausea associated with chemotherapy. Another indication for THC is the treatment of amyotrophic lateral sclerosis (ALS), i.e. potential delay in the progression of the disease and alleviation of the ALS symptoms. CBD on the other hand is a non-psychoactive, anti-inflammatory medicine. It is preferred over THC by patients in need of treatment for arthritis, mood disorders, and nausea.

Studies have shown that THC and CBD have oral bioavailability of approximately 3% to 8% and this varies by formulation and type of vehicles used. This poor bioavailability is in part attributed to the poor gastrointestinal (GI) solubility of these compounds resulting in erratic and variable absorption from the GI tract. Additionally, a significant first pass metabolism may ensue in the liver following GI absorption. These properties render cannabinoids as excellent candidates for Lipid-Based Drug Delivery (LBDD).
Lipid formulations are shown to improve the bioavailability of cannabinoids by improving solubility, intestinal permeability, and enhancement of the lymphatic route of absorption. Among the key mechanisms associated with oral lipid formulations in enhancing bioavailability is the **improved drug wettability and dispersion in the GI fluids, through natural digestion mechanisms that help maintain drug solubilization in vivo**. The effect lasts long enough for the drug to approach the enterocytes for absorption. Recent studies have found that the bioavailability of THC and CBD from a sesame oil formulation was 2.5-fold and 2.9-fold higher compared to a lipid-free formulation (Zgair, 2016, 2017).

However, not all lipids are equal in the nature of their impact on permeability and absorption. Lipid systems may vary by fatty acid chain length and the type/degree of esterification. The choice of excipient is guided by the overall objectives of the dose and the type of dosage form e.g. tablet, capsule, or powder. The formulation may be fine-tuned to improve lymphatic absorption and/or rate of drug release.

Due to their unique properties and versatility, lipid formulations may be designed to tackle several of the below listed bioavailability objectives in a single lipid formulation:

- ✓ Solubilization of the API in the dose
- ✓ Drug dispersion and dissolution in the GI tract
- ✓ Enhanced permeability across enterocytes and wider absorption window
- ✓ Enhanced lymphatic transport of lipophilic drugs

### Excipients for Oral Delivery

Gattefossé offers a range of excipients for achieving one or more of the objectives listed above. They may be categorized by physical state (liquid vs. solid), melting point, HLB value, as well as biopharmaceutical aspects.

As summarized in Table 2, products like **Maisine® CC**, **Peceol**, **Labrafil® M 1944 CS**, **Labrafil® M 2125 CS** and the **Highly Purified Oils** are suitable for oily formulations which may be filled into capsules. They may also be used as oil phase in self-emulsifying formulations. Long chain fatty acids (LFCA) constitute the bulk of the fatty acids found in the products listed, exceptions being **Labrafac®** series, **Lauroglycol™** series, **Capryol™** series, and **Labrasol®** which are composed of medium chain fatty acids (MFCA) with carbon chain lengths of C8 and C10.
Table 2. Excipients by function, category, and suitability for oral dosage forms

<table>
<thead>
<tr>
<th>Excipient Functionality</th>
<th>Applications and Dosage Forms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Highly purified oils</td>
<td>Soybean Oil, Olive Oil, Sesame Oil</td>
</tr>
<tr>
<td>Oily vehicles</td>
<td>Labrafac® Lipophile WL 1349</td>
</tr>
<tr>
<td>Solubilizers/co-surfactants</td>
<td>Geloil™ SC, Geleol™, Maisine® CC, Peceol®</td>
</tr>
<tr>
<td>Liquid and semi-solid surfactants</td>
<td>Lauroglycol™ 90, Lauroglycol™ FCC, Capryol™ 90, Capryol™ PGMC, Plurol® Oleique CC497</td>
</tr>
<tr>
<td>Solid surfactants</td>
<td>Labrafil® M 1944 CS, Labrafil® M 2125 CS, Labrafil® M 2130 CS, Gelucire® 44/14, Labrasol® ALF</td>
</tr>
<tr>
<td>Taste masking, sustained release agents</td>
<td>Gelucire® 48/16, Gelucire® 50/13</td>
</tr>
<tr>
<td></td>
<td>Compritol® 888 ATO, Precriol® ATO 5</td>
</tr>
</tbody>
</table>

Lymphatic vs. Hepatic Absorption

The fatty acid chain length plays a key role in the emulsification, permeation, and route of absorption. The medium chain esters are known for rapid, hepatic absorption, whereas products consisting of unsaturated LCFA tend to stimulate chylomicron secretion and increase lymphatic uptake. The process can enhance the bioavailability of highly lipophilic drugs through preferential absorption within the lymphatic transport system, consequently decreasing first-pass metabolism of the drug in the liver (Hauss, 1998; Charman, 1991; O’Driscoll, 2002).

Since absorption by the lymph means bypassing the liver, formulation with unsaturated LCFA can be a promising strategy for cannabinoids, which are metabolized extensively in the liver. Moreover, lipid excipients may be selected for their surfactive and often self-emulsification properties. Self-emulsification refers to the unique property of certain lipids that disperse spontaneously in aqueous media, forming micellar and lamellar self-assemblies, which are key in maintaining solubility and improving permeability of poorly soluble drugs such as cannabinoids.
SEDDS/SMEDDS

With careful selection of the excipients having the right fatty acid chain length, melting point, and emulsification properties, it is possible to design sophisticated dosage forms. As an example, self (micro) emulsifying drug delivery systems known as SEDDS and SMEDDS are a suitable solution to improving oral drug delivery of poorly soluble and low permeability drugs.

Labrasol®, Gelucire® 44/14, Gelucire® 48/16, Gelucire® 50/13 and the Labrafil® series (Table 2) consist of self-emulsifying excipients that may be in SEDDS / SMEDDS. Several of these products listed are liquid or semi-solid, hence suitable for soft or hard shell capsules.

Other products, such as Gelucire® 48/16 and Gelucire® 50/13 are solids (higher melting points) and so are suitable also for preparation of granules and powders which can subsequently be filled into hard shell capsules or sachets. Alternatively, the powders may be compressed into tablets.

Generally, self-emulsifying formulations provide a rapid (immediate release) profile (Figure 2). Compritol® 888 ATO and Precirol® ATO 5 on the other hand are applicable to taste masking and the preparation of sustained release delivery systems in the form of granules or tablets.

Formulation Guidelines

Further practical information regarding formulation considerations for lipid formulations can be found in our Formulation Guidelines: Developing Lipid Based Formulations for Oral Bioavailability Enhancement.

Additionally, it is possible to develop unique combinations of excipients to simultaneously achieve enhanced bioavailability and sustained delivery. Additional information on sustained release development is available in our Sustained Release Formulation Guidelines.
Skin as Route of Administration

With an average thickness of 1.2 mm and an acidic pH ranging from 4.2 to 6.1, the skin makes up roughly 15% of the total weight of an adult. Certain therapies may target the skin, or a local area beneath the skin. Others target the systemic circulation, often for a sustained delivery. Therefore, good knowledge of the skin’s physiology and the proper role of excipients is essential in formulating effective topical treatments.

Dosage Forms

In certain cases, cannabinoids need to be applied locally on the skin for various purposes such as anti-acne, anti-psoriatic, anti-fungal, analgesics and anti-inflammatory applications. These may be in topical dosage forms having various sensorial properties / textures, and a range of viscosities.

Other possibilities include ointments, foams, and gels. The choice of dosage form will largely depend on the intended use conditions such as spreadability over a given surface area of the skin, frequency of use, and the intended cannabinoid concentration to be delivered. Additionally, enhancers may be added to improve the rate of drug penetration and permeation into the dermis. Another important consideration is whether the formulation is to remedy a brief episode or a lingering ailment. Proven safety and absence of irritation potential become essential for chronic treatments.

Excipients for Topical Formulations

An overview of Gattefossé emulsifiers, consistency agents, and solubilizers suitable for topical preparations is provided in Figure 3. With excellent tolerance / safety profiles and long history of use, these are products of choice for developing stable and luxurious textures.

Each product listed below has unique advantages. Emulsifying bases are ready-to-use formulation systems designed to simplify and to drastically reduce cost/time for development and production; Tefose® 63 is a very mild base with precedence of use in vaginal preparations for many decades; Gelot™ 64 can help emulsify high levels of polar oils; Labrasol® and Labrafil® are permeation enhancers; and others.
enhancing emulsifiers that can be incorporated in all formulations. Likewise, **Transcutol® P**, a hydroalcoholic solubilizer and enhancer is suitable for a wide range of dosage forms. Microemulsions on the other hand are unique in their transparency, low viscosity and superior permeation enhancing capacity.

### Table 3. Excipients by function, category, and suitability for topical dosage forms

<table>
<thead>
<tr>
<th>Excipient Functionality</th>
<th>Ointment</th>
<th>Cream</th>
<th>Foam</th>
<th>Microemulsion</th>
<th>Gel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oily Vehicles / Carriers</td>
<td></td>
<td></td>
<td></td>
<td>Labrafac™ WL 1349, Plurol® Oleique CC 497</td>
<td></td>
</tr>
<tr>
<td>Emulsifying bases for sensitive drugs</td>
<td>Tefose® 63, Gelot™ 64, Sedefos™ 75, Tefose® 1500</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Consistency, viscosity agents</td>
<td>Compritol® 888, Geleol™, Monosteol™</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Liquid O/W Surfactants</td>
<td>Labrasol®, Labrafil® M 1944, Labrafil® M 2125</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>W/O Emulsifiers</td>
<td>Plurol® Diisostearique</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Solvent</td>
<td>Transcutol® P</td>
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<td></td>
</tr>
</tbody>
</table>

**Penetration and Permeation Enhancers - Transcutol® P**

A number of excipients such as the **Labrasol®** and **Labrafil®** series, **Plurol® Oleique CC 497**, **Lauroglycol™** series, **Capryol™** series, and **Transcutol® P** are solubilizers as well as skin penetration and permeation enhancers. They are often used in combination to modulate drug delivery to/or across the skin.

**Transcutol® P** is globally known to solubilize and enhance the penetration of many drugs, notably THC and delivery across skin layers. Using quantitative autoradiography methods, an independent study (Fabin, 1991) measured the penetration of THC in rat skin from three different enhancing formulations: neat Transcutol® P, PEG 400, and a 7:3 blend of propylene glycol/ethanol. The neat Transcutol® P formulation showed the highest and the most stable THC distribution profile across the skin layers. The results for 24 hours of application (Figure 3) show that the THC concentration delivered from neat Transcutol® P was significantly higher than from the other systems.
Figure 3. Effect of formulation on the skin distribution of THC from PEG 400, Transcutol® P and propylene glycol:ethanol formulations

**Transcutol® P** is a safe and effective solubilizer of both hydrophilic and lipophilic compounds. It acts as an enhancer by inducing a transient swelling of the hydrophilic regions of the stratum corneum, rendering it permeable similar to fully hydrated skin. By diffusing into the skin layers, Transcutol® P improves the drug solubility in the skin and its partitioning into the deeper hydrophilic regions of the dermis (Osborne, 2018).

More information is provided in our *Dermacare* and *Topical Drug Delivery* brochures, available on request.

Additionally, we offer:

✓ Regulatory, safety, and precedence of use assistance
✓ Enhancing formulation efficacy
✓ Expertise in sensorial properties
A highly effective and yet unexploited approach to difficult drugs is mucosal delivery. The possibilities include nasal, buccal, rectal, and vaginal routes of administration. In addition to circumventing exposure to the GI tract where drug degradation may occur by digestive enzymes, bile sales and/or pH related effects, this approach helps eliminate any first pass (hepatic elimination) risks for drugs like cannabinoids.

**Mucosal Delivery**

Mucosal delivery is a unique alternative for administration of drug to patients with special needs. Examples include children suffering from recurring seizures, patients unable to take oral medication, and those in palliative care who may receive medicine as nasal sprays or suppositories.

Mucosal delivery is the fastest route of delivery, next to injectable / parenteral routes administration. Compared to oral delivery, the benefits for incapacitated patients include:

- Rapid drug absorption and onset of action
- By-passing the hostile GI tract
- No risks of hepatic elimination
- No taste issue or emesis

**Nasal or Buccal Sprays and Tinctures**

Several products listed in Table 1, including Gatetfossé solubilizers such as Lauroglycol™, Labrafac®, Lipophile WL 1349, Labrafil® M 1944 and Capryol™ series may be used alone or in combination, to obtain drug solutions or dispersions in aqueous or microemulsion systems for use as sublingual drops or nasal sprays.

**Rectal and Vaginal Delivery**

Solid lipid excipients, known in pharmacopoeia as “Hard fat” commonly constitute the “base” or the medium for delivery of drugs in suppository (rectal) or pessary (vaginal). In order to meet the formulation needs for different drugs, dose variations, mold types, and manufacturing speeds, Gatetfossé has developed a wide range of bases: namely the Suppocire® product line for rectal and the Ovucire® line of excipients for vaginal routes of administration (Table 4).
The Suppocire® bases consist of safe, well characterized, and globally referenced excipients which are primarily composed of glycerol esters of fatty acids with varying chain length, degree of esterification, hydrophilicity, and rate of crystallization. Within the Suppocire® line of products, there are unique series, each designed to meet a specific set of formulation and manufacturing conditions. As an example the Suppocire® M series are multipurpose vehicles for a wide range of drugs; the Suppocire® N series are applied in high speed manufacturing environments; and Ovucire® are appropriate for the hydrophilic vaginal environments.

Table 4. Rectal and vaginal product series

<table>
<thead>
<tr>
<th>Product Series</th>
<th>Applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Suppocire® M</td>
<td>Multipurpose bases offering a range of melting points and hydroxyl values</td>
</tr>
<tr>
<td>Suppocire® N</td>
<td>Bases with for sensitive actives and high speed manufacturing processes</td>
</tr>
<tr>
<td>Suppocire® P</td>
<td>Amphiphilic bases for poorly soluble drugs</td>
</tr>
<tr>
<td>Ovucire®</td>
<td>Bases optimized for vaginal drug delivery</td>
</tr>
</tbody>
</table>

The drug properties, intended dose, and mode of absorption are among the critical considerations when selecting an appropriate base that will deliver the optimal physiochemical stability, dispersion and release profile.

Having helped develop hundreds of marketed references over a 60-year span, our group has significant and in-depth knowledge to share with the customers.

Advantages of Suppocire® and Ovucire® Series

- Drug delivery in high concentrations
- Faster absorption for a rapid onset of action
- Alternative for niche patient populations
- Bypassing the hostile GI tract

Detailed information and complete list of product series are available upon request.
Charman W & Stella V (1991)  
_Transport of lipophilic molecules by the intestinal lymphatic system._  

Fabin B & Touitou E (1991)  
_Localization of lipophilic molecules penetrating rat skin in vivo by quantitative autoradiography._  

Hauss DJ, et al. (1998)  
_Lipid-based delivery systems for improving the bioavailability and lymphatic transport of a poorly water-soluble LTB4 inhibitor._  

O’Driscoll CM (2002)  
_Lipid-based formulations for intestinal lymphatic delivery._  

Osborne DW & Musakhanian J (2018)  
_Skin Penetration and Permeation Properties of Transcutol®-Neat or Diluted Mixtures._  
AAPS PharmSciTech.

_Cannabidiol bioavailability after nasal and transdermal application: effect of permeation enhancers._  
Drug Dev Ind Pharm 36(9):1088-1097.

Zgair A, et al. (2016)  
_Dietary fats and pharmaceutical lipid excipients increase systemic exposure to orally administered cannabis and cannabis-based medicines._  

_Oral administration of cannabis with lipids leads to high levels of cannabinoids in the intestinal lymphatic system and prominent immunomodulation._  
We can reduce your development time by providing straightforward formulation guidelines for oral, dermal, rectal and vaginal dosage forms as well as access to extensive databases comprising hundreds of validated placebo or model drug formulations. Our application laboratories in France, India, China and the United States are at your service to provide technical support and formulation feasibility assessment.

We have decades of formulation experience with both experimental and model drugs. We are committed to answering your questions on formulation, regulatory, safety, precedence of use, and scale-up issues.

Please contact your local Gattefossé representative

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