Fluorouracil is a pyrimidine analogue that binds irreversibly within the cell to thymidylate synthetase. This prevents the incorporation of uracil into nuclear RNA which destroys abnormal cancer cells. Fluorouracil cream is used for the treatment of malignant and superficial pre-malignant skin lesions. It is most often prescribed for actinic keratoses and in situ squamous cell carcinoma. It is also occasionally used to treat superficial basal cell carcinomas.

Actinic keratoses are skin lesions caused by long-term ultraviolet (UV) light exposure. These lesions have the potential to develop into squamous cell carcinomas if left untreated.

**EXPLAIN HOW TO APPLY CORRECTLY**

Advising patients to wash the area first with water, dry the skin, then apply a small amount of cream to all the affected treatment areas and gently rub in with a fingertip. Then rinse the finger thoroughly with water. Some patients may prefer to use a cotton bud to apply the cream, or to wear a glove. Fluorouracil cream is absorbed through mucous membranes, so it is very important that patients understand that they should avoid contact of the cream with eyes and the inside of the nose. The lips should be avoided unless it is absolutely necessary on this site. Care should be taken when applying in or near skin folds because it is likely to cause irritation in these sites.

Tretinoin cream is sometimes used prior to starting with a course of fluorouracil cream because it can enhance the effect, and reduce the time required for fluorouracil treatment. Tretinoin peels off the top layer of skin, and works best if it is used for two weeks prior to starting fluorouracil. Tretinoin may also be continued after the fluorouracil course has finished to prevent deterioration and reduce signs of sun damage. Skin must be protected from the sun with sunscreen and exposure to the sun avoided where possible.

**AVOID IN PATIENTS WHO HAVE DIHYROPYRIMIDINE DEHYDROGENASE (DPD) ENZYME DEFICIENCY**

Between 3 and 5% of the population may have partial dihyropyrimidine dehydrogenase (DPD) enzyme deficiency. When using topical fluorouracil, these patients can develop life-threatening systemic toxicity including neutropenia, stomatitis, diarrhoea and neurotoxicity. This may present with fever, chills, fatigue, bloody diarrhoea, vomiting and abdominal pain. Patients should be advised to contact their doctor immediately if they develop any of these side-effects.

**WARN PATIENTS ABOUT SIDE EFFECTS**

It is important to warn patients that adverse effects are expected, particularly inflammation. In some studies, up to 97% of patients have reported at least one adverse effect with topical fluorouracil cream ranging from mild to severe, including erythema, dryness, pruritis, rash, pain, erosion, crusting, changes in skin colour and inflammation. Patients should be warned that the sores and crusts and the resultant raw area that may develop are expected and will heal once the treatment has stopped. However, if any of these symptoms persist or worsen, patient should be advised to consult their doctor or pharmacist.

There may be some systemic absorption which may lead to fever, headache, nausea, diarrhoea, and stomatitis.
Although, life threatening adverse events caused by topical fluorouracil are rare, prescribers should be aware of the possibility and patients advised to contact their doctor immediately if they develop any of these side-effects.\textsuperscript{1,4,6,7}

Always reinforce to patients the importance of safe storage with fluorouracil cream. The cream should be kept in a place that is not accessible by children or other people that may use it inadvertently, for example for a rash.\textsuperscript{1,6}

Pregnancy and breastfeeding
Fluorouracil cream should not be used during pregnancy or breastfeeding.\textsuperscript{1,4,6}

\textbf{ADVISE PATIENTS TO AVOID PROLONGED EXPOSURE TO SUNLIGHT DURING TREATMENT}

It is advisable to stay indoors during the middle of the day during treatment with fluorouracil cream. To avoid excessive exposure to sunlight, it may be preferable for patients to use the cream during the winter months. If treated areas are exposed to the sun, the reaction will be more vigorous.\textsuperscript{1,6}

\textbf{EXPLAIN THAT HEALING MAY NOT BE COMPLETE FOR SEVERAL WEEKS AFTER THERAPY HAS STOPPED}

When fluorouracil cream is applied to the treatment areas, it typically produces the following pattern of response:

- Early and severe inflammatory phase (eg erythema),
- Necrotic phase (eg scaling, tenderness, erosion, ulceration, necrosis
- Healing (eg re-epithelialisation).\textsuperscript{4}

Although the cream should not harm healthy skin there may be subclinical lesions on surrounding skin which will have a similar pattern of response to detectable lesions.\textsuperscript{4,5} The cream usually causes a mild to severe stinging or burning sensation during the treatment period, depending on the skin sensitivity, severity of damage and how long the cream has been used for.

In most cases, after 5-10 days of use the sun damaged parts of the treated skin will become red and irritated. As treatment continues (11-14 days), sores and crust may appear. This is because of the destruction of defective skin cells and is an expected part of the treatment. A dressing can usually be used over these areas depending on patient preference.\textsuperscript{1,4} If the lesions are on the face, once treatment with the cream has finished, make-up or concealer may be used, however this may sting.\textsuperscript{4}

In general, lesions on the face and scalp respond faster than on the limbs, trunk and hands.\textsuperscript{4} Inform patients that healing may not be complete until one or two months after therapy has stopped. Treated areas are often redder than normal and may be more sensitive. This will gradually fade over a few weeks to months.\textsuperscript{1} Reassure patients that scarring is not expected although they should be aware that scarring and hyperpigmentation is a known side effect.\textsuperscript{2} An oily emollient and/or a mild topical corticosteroid cream may help to alleviate discomfort or itch, although care is needed when applying to raw areas.\textsuperscript{1,6}

Ideally, it is best to review patients 2-3 weeks after starting treatment to ensure that there is a therapeutic but not an excessive effect of the cream. Further courses of treatment may be used if necessary, but make sure that patients are aware that they must not self-diagnose or re-start therapy without consulting with the prescriber first.\textsuperscript{1}
FLUOROURACIL

Usual treatment response

<table>
<thead>
<tr>
<th>Duration</th>
<th>Usual symptoms</th>
<th>Patient action</th>
</tr>
</thead>
<tbody>
<tr>
<td>First 5-10 days</td>
<td>Redness and irritation</td>
<td>Avoid direct sunlight</td>
</tr>
<tr>
<td>11-14 days</td>
<td>Blistering, peeling, cracking, sores and crusting, may be some discomfort</td>
<td>May use a light dressing to cover lesion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Arrange appointment with doctor</td>
</tr>
<tr>
<td>14-28 days</td>
<td>Redness</td>
<td>May use white soft paraffin, oily emollient or mild topical corticosteroid cream, if necessary</td>
</tr>
<tr>
<td>2-3 months</td>
<td>Fading</td>
<td>May cover with make up if desired</td>
</tr>
</tbody>
</table>

REFERENCES


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We wish to thank Naomi Petterson, Pharmacist at Waitemata District Health Board, for her valuable contribution to this bulletin.

For further information on other high-risk medicines visit our website at: [www.saferx.co.nz](http://www.saferx.co.nz)