

Everyone's Greasing UP, But Should You Rub It In? A Review of Topical Analgesics and Available Evidence in Clinical Trials

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Disclosure

Consultant/Independent Contractor: AxialHealthcare Inc., Advisory Board: Purdue Pharma LP Honoraria: Auburn, Rockpointe

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Learning Objectives

- Discuss the rationale for compounded topical analgesics.
- Review commercially available topical analgesic options.
- Describe the mechanism of action and clinical applications of topical analgesics.
- Analyze where evidence exists for efficacy with topical analgesics
- Evaluate the efficacy of various topical analgesics and their role in chronic pain.

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Pretest Question #1

Capsaicin 8% patch is approved for which indication in Europe but not in the United States?

- A. Postherpetic neuralgia (PHN)
- B. Dynamic mechanical allodynia (DMA)
- C. Peripheral neuropathy (PN)
- D. Diabetic peripheral neuropathy (DPN)

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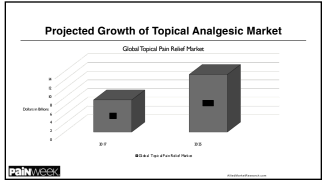
Pretest Question #2

Which prescription oral NSAIDs are also available as prescription topical formulations in the US?

- A. Ketoprofen
- B. Meloxicam
- C. Celecoxib
- D. Diclofenac
- E. All of the above

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Dosage Forms & Delivery Methods

Topical

■Local effect

- Under application site
- •Not intended for systemic absorption Low risk for adverse effects

Transdermal

- •Designed to penetrate into systemic circulation
- Achieve therapeutic plasma concentrations
- •Alternative dosage form
- •Avoid GI or infusion related adverse

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Various Topical Analgesics

| Agent | Availability | Use(s) |
|---------------------------|--------------|--|
| Capsaicin | OTC/RX | -Postherpetic neuralgia |
| | | -HIV neuropathy (off label) -Minor pain |
| Camphor | отс | -Minor pain |
| | 1 | -Pruritus |
| Diclofenac | RX | -Osteoarthritis |
| | | -Acute pain |
| | | -Actinic keratosis |
| Histamine dihydrochloride | отс | -Nociceptive pain relief |
| Lidocaine | OTC/RX | -Postherpetic neuralgia |
| | | -Localized pain |
| | | -Pain and itching of anorectal disorders |
| Menthol | отс | -Nociceptive pain relief |
| Menthol/methyl-salicylate | отс | -Nociceptive pain relief |
| Trolamine salicylate | отс | -Nociceptive pain relief |
| Turpentine | отс | -Nociceptive pain relief |

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Considerations for Topical Analgesics

Advantages

■ Limited Systemic Absorption

- Effective for localized painTissue Concentration > Oral
- Limited Adverse Effect Profile

Disadvantages

- Erratic local absorption
- Variable Depth of Penetration
- Inaccuracy of Dosing
- Require Frequent Applications
 Oleaginous "Greasy" Feeling
 Expensive

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| Salicylate-Containing Rubefacients | |
| Nociceptive Pain | |
| Focus area: | |
| Menthol-methyl salicylate | |
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| Salicylate-containing Rubefacients | |
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| MOA: rubefacients cause irritation of the skin, and are believed to relieve pain in muscles, joints and tendons, and other musculoskeletal pains in the | |
| extremities by counter-irritation | |
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| Irritation of the sensory nerve endings alters or offsets pain in the underlying muscle or joints that are served by the same nerves | |
| muscle or joints that are served by the same herves | |
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| Moore RA, Deny S, McQuay NJ. Cochrane Dalabase Syst Rev. 2010;(7) | |
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| Monthal/mathyl policylete | |
| Menthol/methyl-salicylate | |
| • Menthol is an alcohol (peppermint oil) | - |
| -Topically acts to dilate blood vessels, causing a cooling sensation and analgesic effect | |
| • Methyl salicylate is an ester oil (wintergreen oil) | |
| -Topically induces skin redness and irritation leading to analgesic effect | |
| -Converted to salicylate in the skin | |
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| Derry S. et al. Cochrane Database Syst Rev. 2014;111:CD007403 | 1 |

| Methyl-salicylate – Key Considerations | |
|---|---|
| Methyl salicylate is used as a flavoring agent (inactive ingredient) in oral of products up to a maximum potency of 16mg | drug ———————————————————————————————————— |
| | |
| Allowed as an inactive ingredient in topical gels up to a maximum concentration of 1% | |
| • The maximum systemic salicylate level, in a trial evaluating co-administra | |
| of 10 patches (containing 105mg methyl salicylate/patch) was 0.6782 mg/ -18-fold lower than the minimum value associated with mild toxicity symptoms 20% of topically-applied methyl salicylate may be absorbed | dL |
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| 47 FR 34446 at 34450; December 3, 1992 Center for Drug Evaluation and Research. Application 02-029. Medical Review(s). | |
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| Salicylate-containing Rubefacients- Evidence | <u> </u> |
| Cochrane Meta-Analysis (2014) Acute Conditions, NNT=3.2, RR=1.9 | |
| Chronic conditions, NNT=6.2, RR=1.6 Limitation: quality, validity, and size of available studies | |
| - Limitation. quality, valuity, and size of available studies | |
| Evidence does not support the use of topical salicylate-containing rubefacients for either acute or chronic musculoskeletal pain | 9 |
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| Capsaicin | |
| Nociceptive pain & Neuropathic pain | <u> </u> |
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- MOA: Capsaicin, the pungent component of hot chili pepper, is a vanilloid receptor (VR1) agonist
 - -specifically classified as an agonist of the transient receptor potential vanilloid 1 (TRPV1) receptor
- •TRPV1 is an ion channel-receptor complex expressed on nociceptive nerve fibers in the skin that detect noxious painful stimuli
- Capsaicin causes an initial enhanced stimulation of the TRPV1
- -Depletion of substance P and desensitization
- Analgesia is mediated by death of distal nerve twigs (C fibers)
- -Reversible loss of autonomic & sensory nerve fibers
- -Autonomic nerves recover in 40-50 days, sensory nerves in 140-150 days

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Gibbons CH et al. Ann Neurol. 2010; 68(6):888-898.

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Capsaicin OTC products

Dosage Forms

- Creams (0.025%, 0.075%, 0.1%)
- Patches (0.025%)
- Liquid (0.1%, 0.15%)

Application Tips:

- •Use gloves; wash hands with soap and water after use
- ■Do not use immediately <u>BEFORE</u> or <u>AFTER</u> a bath or shower
- ■Do not use on wounds or damaged skin, with a heating pad, with other external analgesic products

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Capsaicin 8% Patch

- FDA approved for the management of neuropathic pain associated with postherpetic neuralgia (PHN)
- European Medicines Agency (EMA) approved for peripheral neuropathic pain
- Patch (14 cm x 20 cm) 179 mg of capsaicin
- Only physicians or health care professionals under close physician supervision may administer
- -1-4 patch(es) applied for 60-minute duration, frequency not to exceed every 3 months
 -Pre-treatment with topical anesthetic (+/- oral analgesic) prior to application; removal with cleansing gel post-application

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Center for Drug Evaluation and Research. Application 022395. FDA Medical Revie

| Capsaicin- Key Considerations | |
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| Low Dose Capsaicin: | |
| Neuropathic conditions NNT=6.4 (4 weeks), 5.7 (8 weeks) | |
| Musculoskeletal conditions NNT=8.1 Neuropathic and musculoskeletal pain, NNH=9.8 | |
| Todiopalilo dila Habodiosiolotta pali, Hill =0.0 | |
| High Dose Capsaicin: | |
| ■ Neuropathic conditions NNT=6-9 | |
| Adverse events from capsaicin are mainly at the application site (burning, | |
| stinging, erythema) | |
| Painweek, Mason L. et. al. 1811. 2004-328/3744/3901. Deny S. et al. Cochrane Syst Rev 2012 (2): | |
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| Capsaicin-Guidelines |] |
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| American Academy of Neurology (AAN 2004/2011) ■ 2 nd line postherpetic neuralgia | |
| ■ Level B evidence for painful diabetic neuropathy | |
| European Federation of Neurological Societies (EFNS 2010) | |
| 2nd/3rd line for postherpetic neuralgia Level A (8% patch), Level B (cream) efficacy rating for PHN | |
| International Association for the Study of Pain (IASP 2015) | |
| Capsaicin 8% patch, 2nd line for peripheral neuropathic pain syndromes | |
| National Institute for Clinical Excellence (NICE 2017) Capsaicin reasonable alternative to oral medications for peripheral neuropathy | |
| – Oral medications 1 st line | |
| Capsaicin cream > Capsaicin 8% patch | |
| Dubinsky RM, et al. Neurology 2004; 25: 558-55. Ed V, et al. Neurology 2017; 70: 1728-65. Finency NR, et al. Laccet Neurol 2015; 12: 772. Finency NR, et al. Laccet Neurol 2015; 12: 773. | |
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| Lidosoino | |
| Lidocaine Neuropathic pain | |
| Focus Area: Lidocaine 5% patch | |
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| Topical Lidocaine | | - | |
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| MOA: lidocaine is an amide-type local anesthetic agent and is su stabilize neuronal membranes by inhibiting the ionic fluxes requir | | | |
| initiation and conduction of impulses | | | |
| Reduces the frequency rather than the duration of sodium chan | nel opening | | |
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| PainWeek. | J:CD010958. | | |
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| Lidocaine 5% Patch | | | |
| ■FDA approved indication: relief of pain associated with posthe | erpetic | | |
| neuralgia | | | |
| Apply up to 3 patches to most painful areas for up to 12 hours w period | ithin a 24 hour | | |
| -12 hours on/12 hours off | | | |
| -patch is 10 cm x 14 cm containing 700 mg of lidocaine | | | |
| ■ Patches may be cut into smaller sizes prior to removal of the rel | | | |
| Approximately 3 ± 2% of the dose applied is expected to be abs At least 95% (665 mg) of lidocaine will remain in a used patch | orbed | | |
| -At least 95% (665 mg) of ildocaline will remain in a used patch -May be utilized for alternative pain sites | | | |
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| Lidocaine-Key Considerations | | | |
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| Topically administered lidocaine is approximately 70% bound | d to plasma | | |
| proteins - Systemic concentration does not increase with daily use | | | |
| -, | | | |
| Mean peak blood concentration of lidocaine ~0.13 μg/mL | | | |
| ■~1/10 of the therapeutic concentration required to treat cardiac a | arrhythmias | | |
| ■~1/50 of concentrations associated with toxicity (5 μg/mL) | | | |
| –Concentrations higher than 0.25 μg/mL have been observed in some in | ndividuals | | |
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| Topical Lidocaine- Guidelines | |
|---|---|
| American Academy of Neurology (AAN 2004/2011) | |
| ■ 1st line postherpetic neuralgia | |
| ■ 2 nd line painful diabetic neuropathy (Level C Evidence) | |
| European Federation of Neurological Societies (EFNS 2010) | |
| ■ 1 st line for postherpetic neuralgia | |
| | |
| International Association for the Study of Pain (IASP 2015) | |
| 2nd line for mixed neuropathies | |
| National Institute for Clinical Excellence (NICE) (2017) | |
| ■ Reasonable due to safety | |
| Insufficient evidence for efficacy Dubraky RM, et al. Neurology 2064; 53: 959-45. Bit V., et al. Neurology 2061; 78: 1758-45. Finency RM, et al. Laxed Neurol 2016; 17: 1175-23. Finency RM, et al. Laxed Neurol 2016; 17: 1175-23. | |
| Bril V, et al. Neurology 2011; 76: 1758-55. Finnerup NB, et al. Lancet Neurol 2015; 162-73. | |
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| Topical NSAIDS | |
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| Nociceptive pain | |
| Focus area: topical Diclofenac | |
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| Topical NSAIDS | |
| MOA: reversibly inhibit the enzyme cyclooxygenase (prostaglandin endoperoxide | |
| synthase or COX), mediating production of prostaglandins and thromboxane A2 | |
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| Topical application is based on the ability of NSAIDs to inhibit cox enzymes locally | |
| and peripherally, with minimum systemic uptake. | |
| More effective for smaller joints and superficial tissue due to lack of penetration Tissue concentration (subcutis, muscles, tendons) several times higher than oral | |
| - nesure concernitation (subcutis, muscles, tendons) several times higher than oral | |
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Topical Diclofenac Pharmacokinetics Diclofenac Prescription Dosage Forms Brand Name Strength FDA Labeling (ng/mL) (hr) (ng/hr/mL) Class Effect Warnings? 50mg TID 2270 ± 778 6.5 3890 ± 1710 Topical NSAIDs GI Risk Cardiac Risk 53.8 ± 32 10 807 ± 478 1% 48g/day* Is there enough evidence to support labeling? Gel 3% 2g TID × 6 days 5 ± 5 4.5 ± 8 9 ± 19 Solaraze Patch 1.3% BID x 5 days 1.3 - 8.8 120 96 Topical 1.5% 745.2 ± 19.4 ± 9.3 QID x 7 days

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Addressing NSAID Related Concerns

Evans (1995) Case-control Study

• Concluded topical non-steroidal anti-inflammatory drugs were not significantly associated with upper gastrointestinal bleeding and perforation

Petersen B, Rovati S. (2009) Review

Systemic concentrations unlikely to have COX-1 mediated effects like interfere with platelet aggregation or compromise gastric protection

Simon (2009) Double-Blind, Double-Dummy, Randomized Controlled Trial

- Addition of topical NSAID to oral did not significantly increase adverse effects
- Authors conclude combination preferable to increase in oral NSAIDs

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Evans JM, et al. BMJ. 1995;311(6995):22-6. Simon L et al. Pain. 2009; Petersen B, Rovati S. Clin Drug Invest 2009; 29(1):1-9

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Addressing NSAID Related Concerns

Therapeutic Goods Administration (Australia) Safety Review of Diclofenac (2014)

- Query of EMA's Adverse Drug Reporting System (ADRS) -84 reports of adverse events with topical diclofenac
- -3 events when oral diclofenac excluded 2 reports of liver function test abnormalities

 - 1 report of GI bleed
- Safety Review Conclusion:
- -Risk/benefit for topical diclofenac remains favorable
- -Paucity of evidence of serious systemic side effects with topical diclofenac

| Topical Diclofenac- Key Considerations | |
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| ■Topical formulations produce negligible systemic concentrations¹ | - |
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| In comparison with opioids, injectables, and corticosteroids, topical NSAIDS have the lowest NNT (3) to see a benefit for hip and knee OA | - |
| Opioids and corticosteroids do not improve the function and stiffness nearly as well as topical NSAIDS | |
| | - |
| Valename B. Round S. C. de Conglisson 2000; 20(1):1-0 2 Annu W., Nati G. Mankewitz R et al. Ondersprints and Cartings 2010; 18-478-429. | |
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| Topical NSAIDs-Clinical Practice Guidelines | |
| American College of Rheumatology (2012) First line for hand OA, alternative for knee OA | |
| VA/DoD (2014) • Alternative to first line oral NSAIDS for knee osteoarthritis (OA) | |
| NICE (2014) • First line for knee and hand OA | |
| Osteoarthritis Research Society International (OARSI 2014) 1st Line for knee OA (preferred over oral) | |
| No - Graph Management of Fig. and Cross Observations Working Group, Days of You Albara, Day 2914. National Conflict arring Centure Contract Confliction, Missianal Institute for Final Research Section (A. J. America, 1920). 19214. Relicating 102. 4 at J. America, Confliction, Missianal Institute for Final Research Section (A. J. America, Confliction). 19214. Relicating 102. 4 at J. America, Confliction, Missiana, Confliction, Confl | |
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| Focus area: Ketamine, Clonidine, Prazosin, Gabapentin | |
| Compounded Topical Analgesics | |
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| Topical Ketamine | |
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| Peripheral MOA: | |
| NMDA receptor antagonism Toll-like Receptor 4 (TRL4) inhibition | |
| Compounded Formulations: | |
| Concentrations: 0.5%-20% | |
| Numerous co-analgesic combinations | |
| Plasma Concentration Considerations: Generally topical systemic plasma levels below detection (<20ng/mL) | |
| - IV/IM analgesic plasma concentrations: 100-300 ng/mL | |
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| PainWeek. Repair D. Let at Macro Assessingue 2015 April 3 (1) 440-9. Saryon J. Assesh Assig. 2014;18(1) 170-9 | |
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| Topical Clonidine | |
| Peripheral MOA: | |
| 2015 Cochrane Review: • Number needed to treat for an additional beneficial outcome (NNTB) 8.33, [95% CI: 4.3 - 50] | |
| RR: 1.35, [95% Cl: 1.03 -1.77] Concluded may give partial pain relief for only some people with peripheral diabetic neuropathy | |
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| PainWeek. Witasah Ast al Gustiness Challenges Challenges Flowers 2015. | |
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| Topical Prazosin |] |
| Peripheral MOA: | |
| a1-adrenoceptor antagonist | |
| Drummond, et al 2016 Prazosin hydrochloride 1% cream | |
| Inhibited dynamic allodynia in patients with an adrenergic component to pain Inhibited hyperalgesia to stimulation on limb affected by complex regional pain syndrome (CRPS) but not in non-affected limbs | |
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| | Peripheral MOA: Peripheral inhibitory action on the generation of ectopic Suppress the release of substance P and calcitonin gen Blockade of the peripheral glutamate receptors | |
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| | Hiom S, et al. 2015 Retrospective review of 23 patients 6% w/w gabapentin applied three times per day to the affected 11 achieved a clinically meaningful 30% reduction in pain | site (maximal area 20cm2) x 1 month |
| | Concentration Considerations: • Topical gabapentin 6% gel across porcine skin, estimated oral gabapentin (2-20 µg /ml) | peak plasma gabapentin concentration (0.3μg/ml) vs |
| P | Painweek. | Park HJ, et al. Can J Ansesth. 2010;57(7):664-71. Hiers S, et al. Br J Dermatol. 2015;173(1):300-2. |
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Compounded Topical Agent Considerations

- Toxicity reports
- -Unknown safety and efficacy
- Inconsistent with FDA approved route and/or indication
- -Centrally-acting medications delivered peripherally
- Unknown optimal dosing
- Drug combinations not proven safe or effective
- Variation in drug vehicles
- -Lack of standardization
- ■Cost (\$\$\$)

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Evidence for Compounded Topical Analgesics -2013 - Tricare spent \$259 million in 2013 -2014 – DoD spent \$746 million on compounded medications in 2014 -2015 – Medicare Part D spent ~\$500 million ■ Congress required evidence of compounded topical analgesic efficacy -DoD funded study at Walter Reed -August 2015 to February 2018 -399 participants (> 50% female, 43% active military)

Double-blind, Double-dummy, Randomized placebo-controlled trial
 Instructed to apply cream three times a day
 Keep pain diary

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Builder et al. And blind Med. 2019, 172 2029.

| Evidence for Compounded Topical Analgesics | | | |
|---|--|--|--|
| All participants divided into three groups based on type of localized pain Nociceptive pain – Ketoprofen, baclofen, cyclobenzaprine, lidocaine | | | |
| -Neuronathic pain - Ketamine, gabanentin, clonidine, lidocaine | | | |

- -Mixed pain Ketamine, gabapentin, diclofenac, cyclobenzaprine, lidocaine
- Randomized into two groups
- -Topical Analgesics
- -Placebo cream
- Results published February 2019
- -No statistically significant results for any of the three groups compared to placebo

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Summary

- Topical analgesics play an important role in management of localized pain
- Evidence for 1st line use is growing for some types of pain
- Provides solutions to common treatment challenges for pain patients
- Minimal risk of systemic adverse effects

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Pretest Question #1

Capsaicin 8% patch is approved for which indication in Europe but <u>not</u> in the United States?

- A. Postherpetic neuralgia (PHN)
- B. Dynamic mechanical allodynia (DMA)
- C. Peripheral neuropathy (PN)
- D. Diabetic peripheral neuropathy (DPN)

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| Pretest Question #2 | |
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| Which prescription oral NSAIDs are also available as prescription topical formulations in the US? | |
| prescription topical formulations in the 05: | |
| A. Ketoprofen | |
| B. Meloxicam | |
| C. Celecoxib | |
| D. Diclofenac | |
| E. All of the above | |
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| QUESTIONS? | |
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| Albed Marinet Remarch. Topical Pain Faliel Marinet by Therapartic Class. Global Opportunity Analysis and Industry Forecast, 2014-2025. Published June 2018. Accessed July 8, 2016. Availables at Marine Marinet Industrial Control Con | |
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| Centre for Drug Dissalation and Filesamon, Application (2002-2019-EL CRIVES) patch 3-presented 25-freehyt salocylate (10%) Medical Reviewal). Published Filesamy 20, 2008. Accessed July 8, 2018. Available at: http://www.centre.init.org/accessed/2008/2009/2007-2009. Amand 7, Elley K. Topical capacities for patch reseases are throughout potential and metabolisms of action of the new high-concentration capacities for patch. But A J Assessit. 2019;1(5)(4):460-4600. Brandou A, D. Moor, M. Carre and Cassers, "Trackal canadation selected with design and selected and patch 1998;2(5):461-461. Brandou A, D. Moor, M. Carre and Cassers, "Trackal canadation selected with design and selected patch 1998;2(5):461-461. | |
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| Cusco G, Narmisko T, Email E et al. Signationly of operation PS potch venue on pregistation on operation motivated alloyed in patients with peripheral manageable pain. Ear J Pain 2018, 12:270-756. FDA Loodem E, Loui D, Dugaliff CM, Published 20192. A consensed July 8, 2019. A motivate on the "bullet representation of processing Applications 2019201, and processed July 8, 2019. A motivate of the "bullet representation of processing Applications 2019201, and processed July 8, 2019. A motivate of the "bullet representation of processing Applications 2019201, bullet representation patients (PS) and processed July 8, 2019. A motivation of the "bullet representation of processing Applications of PS possing Applications 2019201, bullet representation patients (PS). | |
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