

Biocompatibility and Surrounding Tissue Response to the Buprenorphine in Polymer Biodegradable Polymer Matrix

The determination of in Polymer biodegradable polymer biocompatibility and other information in this bulletin is based upon reports from veterinarians and the study cited below.

- It is important to recognize that placing a biodegradable material in a biological environment will
 initiate a response to the bodies perceived infringement and activate mechanisms to effect a
 healing reaction.
- While inflammation, wound healing and foreign body reaction are considered parts of the tissue responses to "injury", intensity of the inflammatory response greatly contributes to the biocompatibility and successful administration of a drug delivery system.
- When properly injected into a substantial, unrestricted subcutaneous body region, studies verified that the overall inflammatory response remained low, without scarring, tissue necrosis, or suppuration.
- If however, the in polymer formulation goes intradermally it can disrupt the dermal microvascular bed and has also been shown to cause severe localized necrosis due to improperly administered intramuscular injections.
- The extent and magnitude of inflammatory reactions when these polymers react with muscle tissue has been documented as a granulation tissue type of healing response with the presence of macrophages, fibroblasts and foreign body giant cells. (See Fig. 1)
- While the muscle tissues surrounding the injection site do not show irreversible changes, the soft tissue reactions to these materials in an intramuscular environment leads to formation of a connective tissue capsular lesion with localized, low-level inflammation mainly of granulation of
 Notes Regarding Risk of Feline Injection-Site Sarcomas
 - Wedgewood has received no reports of FISS from Buprenorphine in Polymer injections following administration to over 100,000 feline patients since 2011.
 - 2. Recent studies (including Srivastav et al. in JAVMA, 2012) suggest that risk factors for FISS in cats are related to long term corticosteriod injections and adjuvanted vaccines.



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tissue. (See Fig. 1)

time. (See Fig 1.)

During the course of degradation, the

intramuscular implantation lesions are

tissue and continue to dissipate with

gradually replaced by collagenous



Figure 1. A fibrous capsule forms surrounding a degrading poly(DL-loctide-e-coprelactone) following intraunoscalar tissue involvement. (A-D) shows the process of capsular formation 2-4 weeks after implantation; migration of macrophages toward the biomaterial to reinforce the fibrous capsule; the outer part of the biomaterial leaton stars to fingment and begin capsular dissipation; bulk of biomaterial continues to decrease with time, as the fragments of biomaterial also became smaller; degradation continues until leaton is fully resorbed without incident.

NOTE: These injection site occurrences can take over 4-5 weeks to completely resolve.

SOURCE: Den Dannen, W. E. A., Robinson, P. H., Van Wessel, R., Pennings, A. J., Von Lezawan, M. B. M., & Schekenmad, J. M. (1997). Langterm evaluation of degradation and foreign-body reaction of subastancessly implanted poly (DL-betide-r-capmilations). *Journal of Biomedical Materials Research: An Official Journal of The Society for Biometerials and The Japanene Boelety for Biometerials*, 36(5), 357-346.