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Nichiban Co., Ltd.,  
2-3-3, Sekiguchi, Bunkyo-ku  
Tokyo, Japan 112-8663  
Attn: Takashi Miyamoto

**Subject: FDA Compliance Statement for Nichiban's Protape**

Intertek, on behalf of Nichiban Co, has conducted an FDA regulatory evaluation of their Protape. The intended end-use of the tape is to wrap/bundle produce at room temperature conditions. The regulatory evaluation was based on the following:

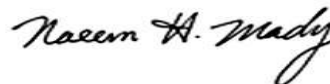
- A comprehensive FDA compliance review of all of the substances used in the manufacture of the Protape
- Toxicological/Safety Assessment conducted on pigments
- Analytical Studies (Ref: RE15506A)

Based on this evaluation, Intertek has determined that Nichiban's Protape is in compliance with US FDA 21 CFR citations for use in food contact applications, intended for use in wrapping/bundling produce at room temperature.

Kind Regards,



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