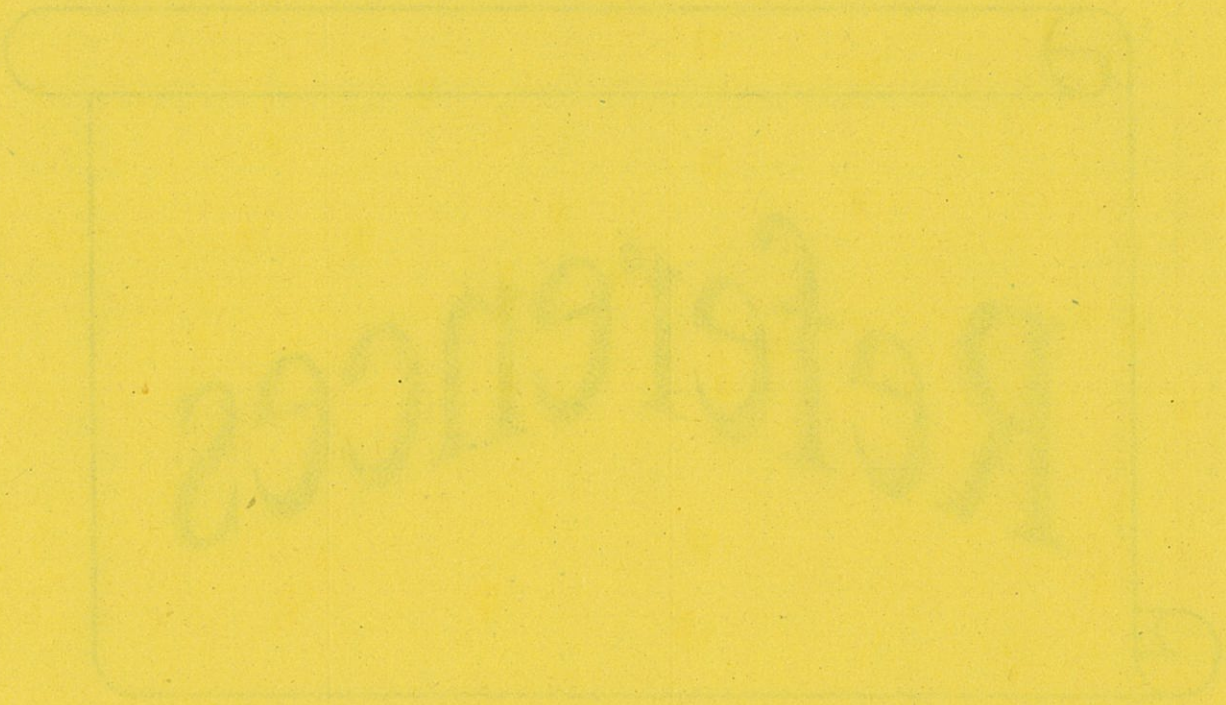


A decorative border on a yellow background, shaped like a scroll. It has a horizontal top bar with a rounded right end and a small loop at the left. A vertical line descends from the left end of the top bar, with a small loop at the bottom. The word "References" is written in a black serif font, tilted upwards from left to right, within the rectangular area defined by the scroll's border.

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