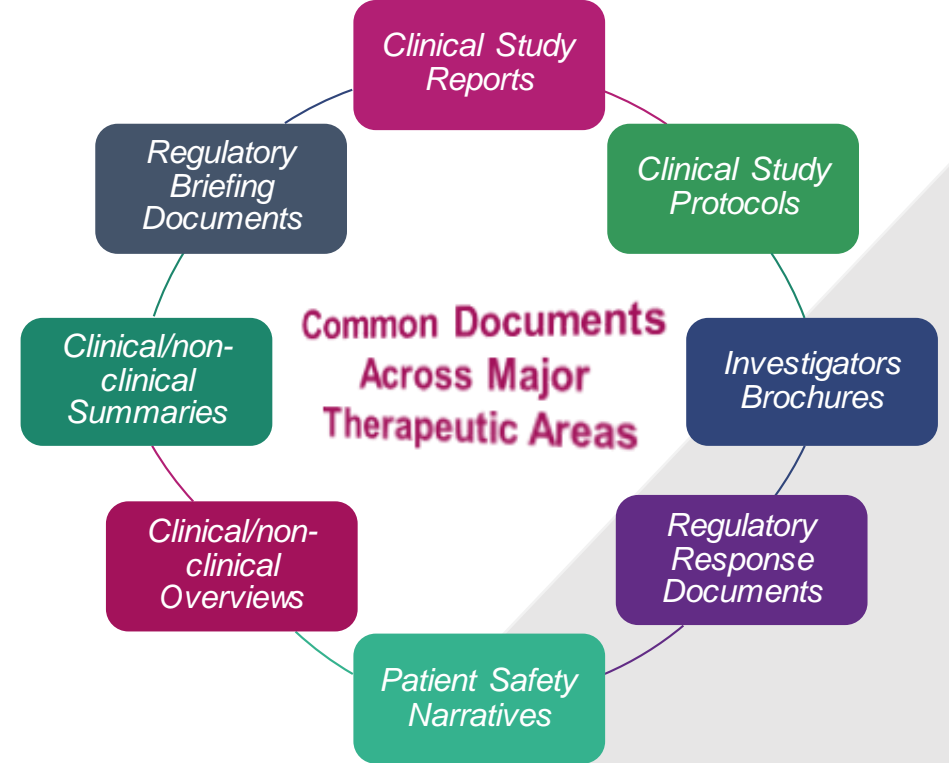


Kinapse Standalone Document QC Service

Poor quality documents pose a risk to patient safety, cause delays in regulatory review and result in wasted effort and budget.

“Consistently achieving Right first time” quality is becoming ever more challenging due to increasing:

- 1 Number of clinical trials
- 2 Amount of regulatory documentation per trial
- 3 Authoring via contractors or vendors outside of the direct control of the sponsor



Many sponsors recognize that independent Quality Control (QC) of documents improves the **quality** and **efficiency** of the regulatory document process.



>9

Years delivering QC services

100%

Success rate in quality and timelines

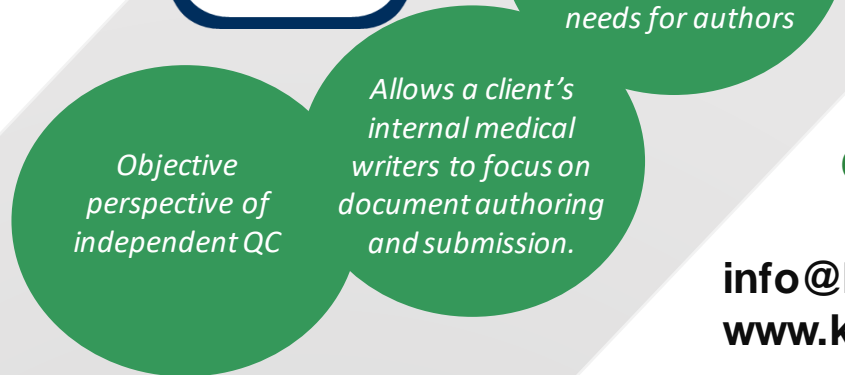


>4000

Documents QC reviewed

4

of top 10 pharma companies are clients



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