

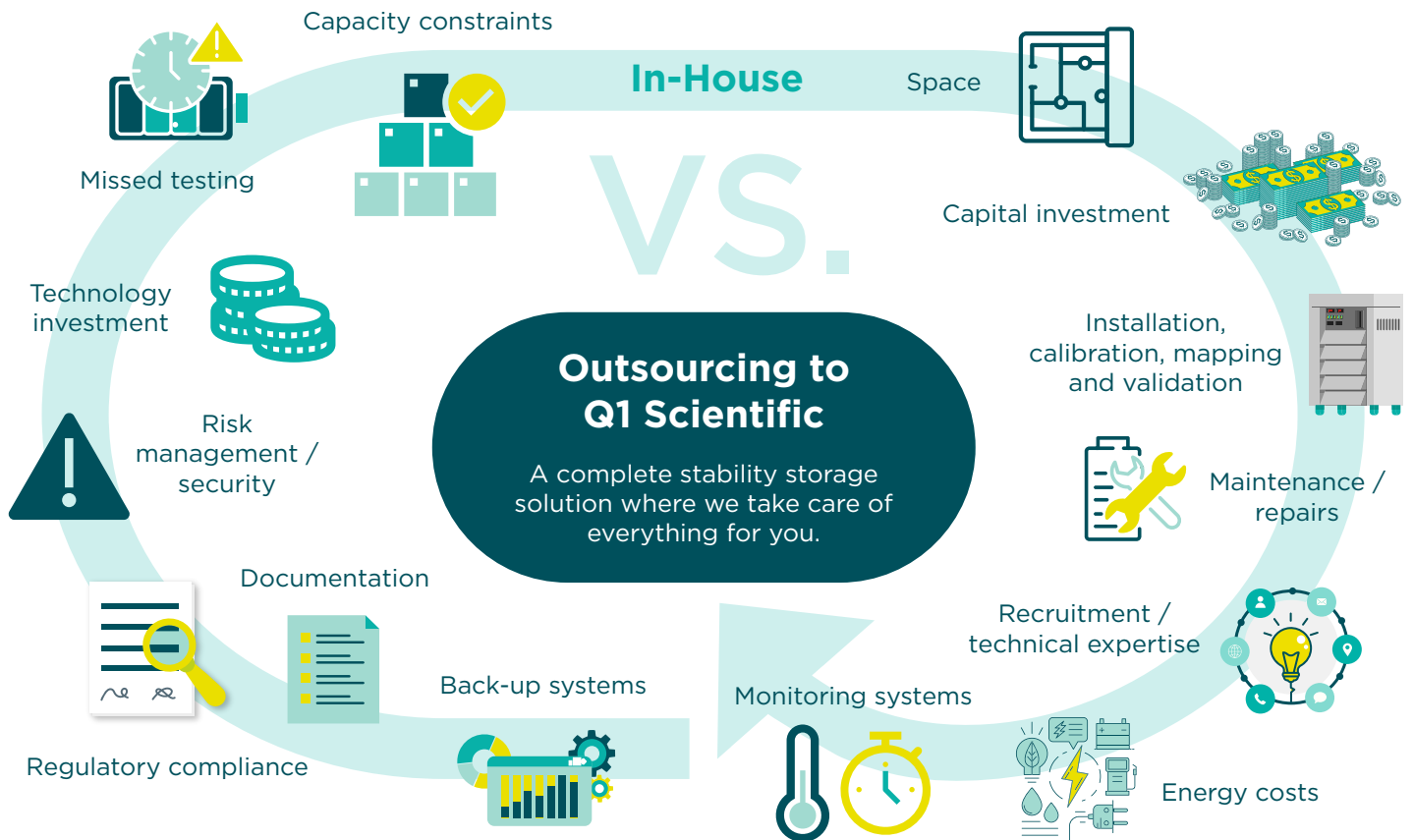
The costs and risks associated with in-house stability storage

Performing stability storage in-house can entail significant costs and risks for pharmaceutical and medical device companies. Some of the costs and risks associated with in-house stability storage are:

Capital expenditure: Purchasing, installing, and validating stability chambers or cabinets requires a large initial investment, as well as ongoing maintenance and calibration costs. Stability chambers or cabinets also occupy valuable space and resources that could be used for other purposes.

Operational expenditure: Operating and managing stability chambers or cabinets incur additional costs, such as electricity, water, personnel, etc. Stability chambers or cabinets also consume energy and generate heat and noise, which can affect the working environment.

Compliance risk: Maintaining and monitoring stability chambers or cabinets poses a compliance risk, as companies need to adhere to the strict regulatory guidelines and standards for stability storage. Stability chambers or cabinets can malfunction or fail due to various reasons, such as power outage, mechanical breakdown or human error, which can compromise the integrity and safety of the stability samples and data.



Outsourcing stability storage to Q1 Scientific can help pharmaceutical and medical device companies save costs and resources, as well as improve their quality, efficiency, and innovation.

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