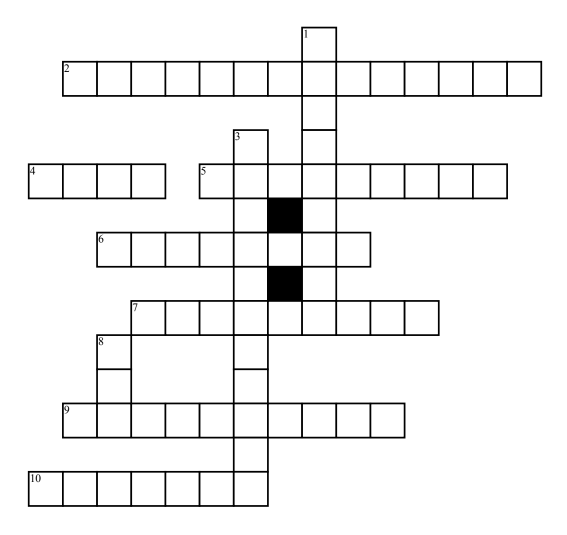
Name:	Date:
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## **GMP** Crossword



## **Across**

- **2.** The nonfulfillment of a specified requirement
- **4.** The combination of the probability of occurrence and severity of harm
- **5.** Regulations apply to all PSI
- **6.** PSI purchases supplies/components from suppliers that have met PSI acceptance
- 7. \_\_\_\_\_ is a preplanned addition, exclusion or deviation from procedure

- **9.** Who is responsible for establishing the PSI Quality Policy/Quality Objectives?
- **10.** All employees are responsible for \_\_\_\_\_

## **Down**

- **1.** Is an allegation of product deficiency received in written, electronic or oral formats
- **3.** When is the best time to notify QA of a non-conformance?
- **8.** Compliance with \_\_\_\_ 21 CFR is required by law

## **Word Bank**

Complaint

Nonconformance Management Deviation Quality Risk Immediately Criteria FDA

**Employees**