

# S.B.SAUDAGAR

Targeting challenging assignment in implementing **Quality Control, Continuous Process Improvements, Customer Relationship Management** and **Team Management** with an organization of repute

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## PROFILE SUMMARY

- ❖ **Strategic & enterprising leader**, with **26 years** of career success in developing & executing operational strategies to accomplish top & bottom-line profitability
- ❖ **Achieving QA operational objectives** by contributing information and analysis to strategic plans and reviews
- ❖ Defined & implemented **production, productivity, quality, and customer-service standards**; identified and resolved problems
- ❖ **Met quality assurance financial objectives** by estimating requirements, preparing an annual budget, scheduling expenditures, analyzing variances, and initiating corrective actions
- ❖ **Developed quality assurance plans** by conducting hazard analyses; identified critical control points and preventive measures
- ❖ Coordinated the Manufacturing **Non-conformance process (NC)** for all Production areas ensuring compliance to Quality and Regulatory Standards
- ❖ **Led cross functional teams** providing quality guidance and support in the determination of appropriate Non-conformance root cause analysis and implementation of effective corrective actions
- ❖ Experience of implementing **Seven Quality Tools, Quality standards, 5S, 8D, kaizen, Kanban, Poka-Yoke, International Standards (IS)** with quality techniques for improving the operational efficiency
- ❖ Proven capabilities in **determining business unit's mission direction, formulated initiatives & provided strategic advice** for achieving corporate strategic goals; skills in formulating strategies aimed at driving business growth

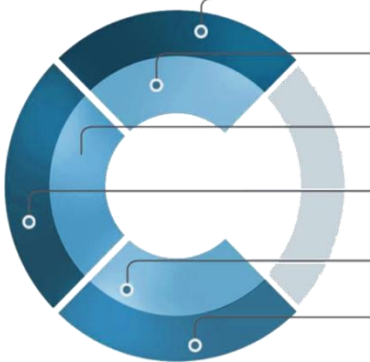


## CORE COMPETENCIES

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|  Quality Assurance & Control |  Risk Assessment & Analysis         |  Profitability Management |
|  Quality Procedures          |  Good Manufacturing Practices (GMP) |  Stakeholder Management   |
|  Reports & Documentation     |  Cross-functional Coordination      |  Team Building & Training |



## SOFT SKILLS



## EDUCATION & CREDENTIALS

- ❖ **M.Sc. in Organic Chemistry** with 63.50% in 1992
- ❖ **B.Sc. in Chemistry** with 56.83% in 1990



**Poona University.**



## MAJOR TRAINING

- ❖ Training Programme on **FTIR Spectroscopy & Gas Chromatography** conducted by Perkin Elmer Instruments Pvt. Limited
- ❖ Proficiency in **Testing & Validation** conducted by Vishwa Bharti Charitable Trust at Mumbai
- ❖ Training programme on **NABL Accreditation Awareness** at IDEMI Mumbai



## WORK EXPERIENCE

**Since Dec-23 with Chetas Biochem-Jaipur [Raj.] Sr.Chemist with Factory Incharge.**

### Key Result Areas:

- ❖ Assuring consistent quality of manufacturing process by developing and enforcing good automated manufacturing practice (GAMP) systems, validating processes, providing documentation, and managing staff
- ❖ Accomplishing quality assurance human resource objectives by recruiting, selecting, orienting, training, assigning, scheduling, coaching, counseling, and disciplining employees and planning and reviewing compensation.
- ❖ Establishing critical limits, monitoring procedures, corrective actions, and verification procedures and monitoring inventories
- ❖ Achieving financial objectives by preparing the quality assurance budget, scheduling expenditures, analyzing variances, and initiating corrective actions
- ❖ Analyzing, modifying and implementing procedures for various departments to ensure that the company's policies and various state regulations were within compliance
- ❖ Identifying process and systems improvements, participating in teams and coordinating ad hoc projects to improve and support order management
- ❖ Identifying deviations and ensuring proper corrective actions by initiating Problem Investigation Forms and utilizing CAPA software in compliance with policies, procedures and regulations
- ❖ Reviewing and approving all quality planning documentation (FMEA, CTQs, SW QA Plan etc.) for new model programs in support of program milestones
- ❖ Actively participating in development and quality reviews, including feedback from customer / system testing teams, to develop and drive quality improvement roadmaps
- ❖ Understanding and maintaining compliance with local and global policies and procedures
- ❖ Communicating quality issues to responsible stakeholders to maintain customer quality requirements
- ❖ Identifying any potential quality issues per defined process and escalating potential quality issues immediately to management
- ❖ Carrying out quality tasks in production with regard to quality complaint
- ❖ Steering efforts in implementation and advance development of the product and process standards required by the QM system

**Oct'20-Oct'21 with Goodway Chemicals Pvt. Ltd., Sarigam as QC-Executive**

### Key Result Areas:

- ❖ Conducted analysis of Finished Products, Raw Material, In-process Quality Control in Pharmaceuticals, Cosmetics, Petrochemical Industries
- ❖ Hands-on with analytical instruments like G.C., HPLC, FTIR, UV-Visible Spectrophotometer and other laboratory related instruments
- ❖ Proven expertise in conducting Classical Chemical Analysis
- ❖ Steered efforts in preparation of Standard Operating Procedure, Analytical Reports, Analytical Method Development Reports
- ❖ Well versed with Good Manufacturing Practices, Good Laboratory Practices, ISO9001 Quality Management System
- ❖ Organized daily qualitative & quantity work in QC & shouldered responsibility for testing of Raw-Materials, Packaging Materials, In-process Materials, and finished goods
- ❖ Played a key role in investigation of specification results, market complaints investigation, long -term stability & monitoring of drugs products, re-testing of raw material and packing material and workload review
- ❖ Leveraged skills in controlling of retention samples of raw materials and finished products
- ❖ Executed stability studies of finished as well as raw materials as per guidelines
- ❖ Learnt various techniques on Characterizing , testing & Evaluation of additives for automotive & Industrial lubrication
- ❖ Extensively used analytical techniques such as GC (NETEL Michro-9100), HPLC (Varian Prostar & Perkin Elmer, Agilent), FTIR (PerkinElmer, Agilent), ICP (Inductively Coupled Plasma-OES/Agilent) Nitrogen Analyzer, Total Acid No. & Total Base No. Evaluation using Potentiometric Titration Auto Titrator (Metler DL-21/25, 50), Kinematic Viscosity Measurement (100 & 40 C) & other Rheological studies on cold temperature properties of oils by CCS, MRV, Pour Point & Brookfield. (Ducom)

**Oct'08-Nov'20 with Century Rayon Ltd., Sahad, Kalyan as Supdt. T.C. Lab**

### Key Result Areas:

- ❖ Organized daily qualitative & quantity work in QC & shouldered responsibility for testing of Raw Materials, Packaging Materials, In-Process Materials, and Finished Goods
- ❖ Steered efforts in investigation of specification results, market complaints investigation, long -term stability & monitoring of drugs products, re-testing of raw material and packing material and workload review
- ❖ Leveraged skills in controlling of retention Samples of raw materials and finished products
- ❖ Executed stability studies of finished as well as raw materials as per guidelines
- ❖ Played a key role in implementation of all quality system, operation and calibration of all laboratory equipment's, ensuring product safety requirements in the laboratory
- ❖ Managed Extraction, Fractionation, Isolation, Purification using various chromatographic techniques and characterization of various groups of drugs compounds
- ❖ Coordinated with various function to resolve quality issues
- ❖ Wealth of expertise in handling of analytical instrument like L.O.D,KF (Karl-Fischer), IR (Agilent) MP,BP, UV Spectrophotometer, PH Meter, Finish Product Assay by "Auto Titrator (Lab India, DL-25,50) & other analytical instrument & various related work study etc.



## PREVIOUS EXPERIENCE

June'07-May'08 with NRC Ltd, Mohone as Sr. Officer-Q.A.

Dec'05-Oct'06 with Shirdi Chemical Pvt. Ltd., Pawane, Navi Mumbai as Analytical-Chemist (Q.C.)

Dec'04-June'05 with Sudarshan Chem. Ind. Ltd. as Sr. Chemist (Analytical Lab)

Apr'04-Nov'04 with Modepro India Pvt. Ltd. as Chemist (Q.C.)

Nov'93-May'03 with M/s. Lubrizol India Pvt. Ltd. Navi Mumbai as Sr. Research-Chemist

❖ Dec. 12-2023 continue with M/s.Chetas Biochem.-Jaipur [Raj.] as Sr.Chemist with Factory Manager.

❖ **July 15th-2024 continue with M/s.Metatech Process Solution -Kharghar -New Bombay as Sr.Chemist with Factory Manager.**

## PERSONAL BRIEF:

- **Name: Sunil Baburao Saudagar.**
- **Father's Name: Baburao S Saudagar.**
- **Date of Birth: 15/08/1967.**
- **Nationality: Indian.**
- **Marital Status: Married.**
- **Address: Hari Vihar.A-607.Near Bacchav Classes. Jail Road.**

**Nashik Road.Nashik-422101 [M.S]**

**"The Above all information is true to the best of my knowledge & Brief".**

**Yours Faithfully**

**[Sunil Saudagar]**

**Place:Panvel-New Bombay**

**Date:15/11/2025**

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**From:**

**S.B.SAUDAGAR**

Hari vihar, A-607.Near

Bacchav Classes.Jail Road, Nasik road, Nasik -422

101. [Maharashtra]

**Contact: 09930665853.**

**Email id:sursun2811@gmail.com**

**To,**

**Personnel Manager [H.R. Dept.]**

**Respected Sir,**

**Subject: Application for the post of “Quality Control/R & D”.**

With reference to your advertisement published in -----dated  
on----- for the post of QC-Executive /Officer/Asst. Manager, I would  
like to apply for the same. My CV and relevant certificates are enclosed for kind

perusal.

**Thanking you.**

**Yours Faithfully.**

**[S. B. Saudagar]**