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| **Mohammad Manaf Aljijeny – 2008074**  To interview this candidate, please send your company name, vacancy, and salary offered details along with this or other CV Reference Numbers that you may have short listed from <http://www.gulfjobseeker.com/employer/cvdatabasepaid.php>  addressing to HR Consultant on email: [cvcontacts@gulfjobseekers.com](mailto:cvcontacts@gulfjobseekers.com)  We will contact the candidate first to ensure their availability for your job  and send you the quotation for our HR Consulting Fees. |



Personal Information

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| |  |  |  | | --- | --- | --- | | **Birth Date:** | 18 november 1983 |  | | **Gender:** | Male | | **Nationality:** | Jordan | | **Residence Country:** | Ajman, United Arab Emirates | | **Visa Status:** | Residency Visa | | **Name in Arabic:** | محمد الججيني | | **Marital Status:** | Married | | **Driving License Issued From:** | Jordan | |



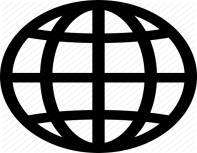
Experience

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| |  |  |  | | --- | --- | --- | | January 2016 - September 2016  **Senior Supervisor** | | | | **Location:** | Amman, Jordan | | | **Company Industry:** | Pharmaceutical | | | **Job Role:** | Quality Control | | | **Key Responsibilities:**   * Manage QC Lab, equipment and data systems. * Manage QC Stability office, stability studies, stability requirements (new products, Annual commitment, Process validation … etc.). * Create, revise, review and approve protocols, supporting analytical data, and reports associated with method validation, testing, and release of GMP starting materials, intermediates, and APIs. * Review and approve Certificate of Analysis, Certificate of Test, and stability studies. * Serve as QC lead on analytical projects teams and product operational teams. * Collaborate with regulatory to support ANDA filling/amendments. * Train direct reports on QC job related functions and technical methods. * Approved and review quality records, processes and documents, including method validation, CAPAs, Change Controls and SOPs. * Ensure that procedures and specifications are appropriate and followed. * Approve and review Transfer of Analytical procedures (TAP). * Distribute laboratory testing of in-process and drug product using validated methods against scientifically-derived, fit-for-purpose specifications. * Approve or reject drug products manufactured, processed, packed. * Ensure investigation of nonconformance (Out of Specification (OOS) / Out of Trend (OOT)). * Ensure investigation is conducted and root cause is eliminated for production and control record errors, discrepancies, and failure to meet specification, including quality attributes. * Review complaints to determine if it relates to a failure to meet specification, if so investigate and report. | | | | | January 2012 - January 2016  **Supervisor**  at Hikma Pharmaceutical Company | | | | | **Location:** | | Amman, Jordan | | | **Company Industry:** | | Pharmaceutical | | | **Job Role:** | | Quality Control | | | **Key Responsibilities:**   * Manage QC Lab, equipment and data systems. * Review and approve Certificate of Analysis, Certificate of Test, and stability studies. * Train direct reports on QC job related functions and technical methods. * Ensure that procedures and specifications are appropriate and followed. * Distribute laboratory testing of in-process and drug product using validated methods against scientifically-derived, fit-for-purpose specifications. * Ensure investigation is conducted and root cause is eliminated for production and control record errors, discrepancies, and failure to meet specification, including quality attributes. | | | | |  | | | | | January 2011 - January 2012  **Senior "Data review"**  at Hikma Pharmaceutical Company | | | | | **Location:** | | Amman, Jordan | | | **Company Industry:** | | Pharmaceutical | | | **Job Role:** | | Quality Control | | | May 2007–December 2010  **Senior Stability Officer**  at Hikma Pharmaceutical Company | | | | | **Location:** | | Amman, Jordan | | | **Company Industry:** | | Pharmaceutical | | | **Job Role:** | | Quality Control | | | |  |  | | --- | --- | | September 2005 - April 2007  **Analyst**  at Hikma Pharmaceutical Company | | | **Location:** | Amman, Jordan | | **Company Industry:** | Pharmaceutical | | **Job Role:** | Quality Control | | | | | |



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| |  |  |  |  |  |  |  |  |  | | --- | --- | --- | --- | --- | --- | --- | --- | --- | | |  |  | | --- | --- | | **Bachelor's degree , Chemistry**  at Al albyat university | | | **Location:** | Mafraq, Jordan | | **Completion Date:** | June - 2005 | | **Grade:** | 79 out of 100 | | |

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| Sk   |  | | --- | | **HPLC/GC**  **Level:** Expert | | **Quality Control Laboratory**  **Level:** Expert | | **Calibration**  **Level:** Expert | | **Microsoft Office**  **Level:** Intermediate | | **LeanSixsigma (Green Belt)**  **Level:** Beginner | |



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| |  | | --- | | **Arabic**  **Level:** Native | | **English**  **Level:** Intermediate | |

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