

BAB 5

KESIMPULAN

Berdasarkan hasil kegiatan Praktek Kerja Profesi Apoteker (PKPA) di Industri Farmasi yang dilakukan secara daring melalui proses pembelajaran dengan sistem diskusi dan sidang pleno yang dilaksanakan pada tanggal 2 Agustus 2021 hingga 25 September 2021 dapat disimpulkan bahwa:

1. Apoteker memiliki peran penting dalam suatu industri farmasi untuk menjamin keamanan, kualitas, dan efikasi sediaan farmasi yang diproduksi.
2. Mahasiswa calon apoteker dapat lebih memahami peran, fungsi, posisi, dan tanggung jawab apoteker dalam industri farmasi.
3. Mahasiswa calon apoteker mendapat wawasan, pengetahuan, keterampilan, dan pengalaman praktis untuk melakukan pekerjaan kefarmasian di industri farmasi, serta lebih mempersiapkan diri agar mampu memasuki dunia kerja sebagai tenaga farmasi yang profesional.
4. Mahasiswa calon apoteker dapat mempelajari prinsip CPOB dan penerapannya dalam industri farmasi.
5. Mahasiswa calon apoteker mendapat gambaran nyata tentang permasalahan pekerjaan kefarmasian di industri farmasi meskipun kegiatan dilakukan secara daring.

BAB 6

SARAN

Saran yang dapat disampaikan dari hasil pelaksanaan kegiatan Praktek Kerja Profesi Apoteker (PKPA) di Industri Farmasi secara daring yang dilaksanakan pada tanggal 02 Agustus 2021 hingga 25 September 2021, sebagai berikut:

1. Pihak praktisi dari masing-masing industri diharapkan terus meningkatkan kerja sama dengan program studi profesi apoteker dari berbagai perguruan tinggi dalam pelaksanaan praktek kerja profesi apoteker.
2. Sebelum melaksanakan kegiatan PKPA di industri, mahasiswa calon apoteker diharapkan lebih banyak membekali diri dengan ilmu pengetahuan tentang kegiatan-kegiatan, perundang-undangan farmasi dan segala hal yang berkaitan dengan industri farmasi.
3. Mahasiswa calon apoteker diharapkan lebih aktif dan tanggap selama menjalankan praktek kerja profesi agar para calon apoteker mendapatkan pengetahuan yang lebih banyak sehingga mampu mencapai semua apa yang menjadi tujuan yang direncanakan.
4. Fasilitator dapat memberikan gambaran dari industri farmasi kepada mahasiswa melalui gambar atau video ataupun media digital lain agar mahasiswa PKPA secara daring dapat menerima gambaran praktek sesungguhnya di lapangan dengan lebih jelas.

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