



علوم شیمی و نفت / شیمی تجزیه و شیمی فیزیک

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کتاب

■ کاربرد نظریه ی گروه در شیمی

ناصر صفری، منصور زاهدی

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پایان نامه ها و رساله های دکتری

بررسی مکانیسم تخریب هیم و تبدیل آن به بیلی وردین و وردوهیم و مطالعات اثرات آنزیم – پروتئین با استفاده از شبیه سازی دینامیک مولکولی

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