



علوم شیمی و نفت / شیمی تجزیه و شیمی فیزیک

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علاقه پژوهشی

■ شیمی محاسباتی

■ طیف سنجی مولکولی

فعالیت‌های اجرایی

■ مدیر گروه آموزشی شیمی محض، ۱۳۹۳-۱۳۹۴

■ استاد راهنما، ۱۳۸۹- تا زمان حال

کتاب

■ کاربرد نظریه ی گروه در شیمی

ناصر صفری، منصور زاهدی

دانشگاه شهید بهشتی - تهران، ایران، ۱۳۹۵، شابک: ۹۷۸۹۶۴۴۵۷۳۵۵۲

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ناصر صفری، منصور زاهدی

انتشارات پیام مولف، ایران، ۱۳۸۳، شابک: ۹۶۴۷۹۱۹۱۹۰

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بررسی نظری مکانیسم کاتالیستی و مهار آنتی بیوتیک های بتالاکتام بامتالوبتالاکتامها در حلال ها و دماهای مختلف با استفاده از محاسبات

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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

فاطمه السادات علوی

۱۳۹۷

■ مطالعه ساختار الکترونی برخی از گونه های اتمی و مولکولی لیتیم و بریلیم با استفاده از روش مونت کارلوی کوانتومی

سعید نصیری

۱۳۹۶

■ بررسی امکان باز تولید ویژگی های مولکولی بر اساس خصوصیات اتم های آزاد: چگالی الکترونی و انرژی همبستگی

زهرا علی محمدی کیوانی

۱۳۹۵

■ . مطالعه نظری خاصیت آنتی اکسیدانی ترانس – رزوراترول و مشتقات آن در فاز گازی، آب و بنزن

الیاس نظرپرور نوشادی

۱۳۹۳

■ . بررسی نظری MgO : مطالعات سطوح کاتالیستی طلا و بر همکنش آن با مولکول اکسیژن و سطح

علی مقدسی

۱۳۹۲

■ مطالعه نظری بر روی جنبه های ساختار لیگند دارویی پامیدرونات و رادیو داروی پامیدرونات – ساماریوم بر اساس روش های شیمی کوانتومی

مسعود عربیه

۱۳۹۱

■ مطالعات نظری فرایند گلایکه شدن با استفاده از روش های شبیه سازی دینامیک مولکولی و شیمی کوانتومی

رسول نصیری

۱۳۹۰

پایان نامه های کارشناسی ارشد

■ . DFT بررسی رنگ خوراکی ایندیگوکارمین و مقایسه آن با سایر رنگ های خوراکی مجاز با استفاده از روش محاسباتی

احمد بختیاری زاده

۱۴۰۰

■ مطالعه نظری طیف های کلاسترهای لیتیم

ناصر واعظی

۱۳۹۸

■ با بروفن خالص و تغییر یافته HF مطالعه نظری بر هم کنش بین گاز

■ مطالعات سطوح کاتالیستی طلا و برهم کنش آن با مولکول هیدروژن بررسی نظری

افشین چراغی

۱۳۹۸

■ بررسی نظری خوشه های آنیونی هیدروژن

امین محمدی

۱۳۹۶

■ نانو ساختارهای سیلیکون کاربیدی: مطالعه نظری بر روی نحوه تشکیل و ویژگی ها و نقایص ساختاری

تاده ظهرا بیان سروشکانی

۱۳۹۵

■ مطالعه نظری ترکیبات اکسیم و مشتقات آن جهت فعال سازی آنزیم استیل کولین استراز

سمانه خواجه وند

۱۳۹۵

■ مطالعات نظری ساختارهای الکترونی نانوگرافن

زینب محبی نیا

۱۳۹۵

■ برای ذخیره سازی هیدروژن $(X_2O_6)(SiH)(X; B, Al, Ga, N, P, As)$ مطالعه نظری خواص هترو فولرن های

میلاد بهرامی فیل ابادی

۱۳۹۴

■ . مطالعه داکینگ بر روی برهمکنش های مشتقات تاکسول با پروتئین بتا – توبولین به منظور معرفی داروی ضد سرطان کارآمدتر

وحیده پورقاسم

۱۳۹۳

■ . بررسی تئوری مسیر اکسید و احیای کبالت اکسوفلورین و اثر لیگند اکسیال بر آن

آذر استوان

۱۳۹۲

■ . فعالیت کاتالیستی برجسته نانو کلاسترهای طلا و آلیاژهای آن : مطالعه تئوری

فاطمه مخلوقی

۱۳۹۲

■ با استفاده از نظریه تابعی دانسیته $[M(=Fe, Co, Ni, Zn)](M(dipic)(H_2O)_3)$ مطالعه نظری ویژگی های ساختاری کمپلکس های

زهرا آقامیرزائی

۱۳۹۱

. DFT به روش MgO مطالعات نظری بر روی انرژی جذب نانو کلاسترهای طلا روی سطح منیزیم اکسید (۱۰۰) و ارتباط آن با نقص های شبکه ای



سمیرا ناصریان مغانی

۱۳۹۱

مطالعات نظری بر روی توزیع چگالی الکترونی نانو کلاسترهای طلای جذب شده روی سطح منیزیم اکسید (۱۰۰) و ارتباط آن با انرژی جذب به

■ DFT روش

زهرا صادقی

۱۳۹۱

■ (بررسی بر همکنش آمینواسیدهای هیستیدین و لایزین با کاتیون منگنز – بررسی کلیه آمینو اسیدها و کاتیون فلزی منگنز) در محیط گازی

حمیدرضا رئیسی هارونی

۱۳۹۰

■ E مطالعه تئوری اثرات استخلاف و حلال بر روی خاصیت آنتی اکسیدانی ویتامین

میثم نجفی

۱۳۹۰

■ مقایسه ساختاری هموگلوبین انسان و غاز با استفاده از شبیه سازی دینامیک مولکولی

ارسلان نیک پسند

۱۳۸۹