among the youngest societies in the developed world. Further efforts are required to reduce the substantial CHF burden.

PHS102
WILL U.S. PAYERS CHAMPION BIOSIMILARS?
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OBJECTIVES: U.S. healthcare spending on high-cost biologics has escalated over the past decade. The Biologics Price Competition and Innovation Act allows for biosimilar products in the United States, which will be lower-cost alternatives to reference biologics. This study explored payer attitudes to the reimbursement and adoption of biosimilars in the United States. METHODS: Some 61 U.S. medical or pharmacy directors at managed care organizations (MCOs) were surveyed during 2014 regarding their expectations for biosimilar reimbursement. RESULTS: Respondents indicate that MCOs expect biosimilars to offer a significant discount to the reference product. However, rapid formulary inclusion of biosimilars is deemed adequate for reimbursement, while 33-34% is deemed necessary for preferential reimbursement to the reference brand. Rapid formulary inclusion of biosimilars is expected, with 79% of respondents indicating formulary inclusion within 12 months of launch. The most conducive uptake strategies are expected for products with deep discounts, to step-therapy requiring biosimilar prescribing prior to the reference brand. Widespread reimbursement of biosimilars in extrapolated indications is, however, uncertain, with only 34% of respondents reporting that their MCO would unconditionally reimburse a biosimilar under such circumstances. Payers expect to employ various strategies to promote biosimilar uptake, from favorable tiering to step-therapy requiring biosimilar prescribing prior to the reference brand. The most conducive uptake strategies are expected for products with deep discounts. In addition, payers will run educational campaigns for physicians. However, the most conducive uptake strategies are expected for products with deep discounts.

PHS103
ARE GENERIC MEDICINES GOOD FOR MY PATIENTS? FINDINGS FROM A QUALITATIVE ASSESSMENT OF PERCEPTIONS AMONG MEDICAL SPECIALISTS IN MALAYSIA
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OBJECTIVES: Medical specialists have an essential role in promotion of generic medicines. However, misconceptions and negative perceptions about generic medicines are prevalent among the public. Therefore, the aim of this study was to explore their knowledge, perceptions and views about generic medicines. METHODS: A qualitative methodology was adopted in this study. A descriptive qualitative (QD) study with phenomenology overtones was conducted. A face-to-face semi-structured interview was conducted with a purposive sample of medical specialists until saturation of data was achieved. The interviews were audio taped and transcribed verbatim. Then, the subsequent transcripts were analyzed using thematic analysis. The themes were generated using both deductive (theoretical) and inductive coding. To ensure reliability of data analysis, peer review and independent analysis of the data was done. RESULTS: Five major themes were identified: (1) factors affecting specialists’ perceptions about generic medicines; (2) specialists’ knowledge and confidence with Malaysian generic regulatory approval system; (4) drug information sources used by the specialists and (5) International non-proprietary name (INN) prescribing. CONCLUSIONS: The study findings showed that several factors influenced prescribing decision with regards to the brands of the medicines including familiarity and experience with generic medicines, medical and clinical condition of the patient, availability of the medicine brand in the hospital and departmental financial allocations. Moreover, misconceptions about quality, safety, efficacy and bioequivalence of generic medicines were prevalent among the participants. Furthermore, the participants were not familiar with the Malaysian generic medicines approval system. Therefore, medical specialists’ concerns need to be addressed. Some recommendations and suggestions were made to improve generic medicines utilization in the country.

PHS104
PROTON PUMP INHIBITOR PRESCRIBING TRENDS IN THE US AMBULATORY SETTING
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OBJECTIVES: The study aimed to examine the utilization patterns of Proton Pump Inhibitors (PPIs) among the elderly and the trends in PPI prescribing across ph ysician specialties, to obtain a wide comprehensive view, and to identify any trends in the utilization of PPIs and their effect on the patient and physician-level factors affecting non-indication based PPI prescription in the above population. METHODS: This retrospective cross-sectional study used data from 2002-2010 National Ambulatory Medical Care Survey (NAMCS) and outpatient visits to office-based physicians. Data from 2005 – 2010 were used to determine data to evaluate PPI utilization patterns based on patient visits. PPI prescribing trend among different physician specialties was estimated using 2005-2010 NAMCS. Evidence-based recommendations for PPIs included the US Food and Drug Administration (FDA) approved indications and the National Institute of Clinical Excellence (NICE) guidelines. Multivariable logistic regression model was performed to assess the association between non-indication based PPI use and various patient and physician-level characteristics using 2008-2010 NAMCS data. RESULTS: The use of PPIs increased from 5.29% patient visits in 2002 to 11.82% in 2010 (p<0.001). Prescriptions of PPI without an appropriate indication by primary care physicians (54.9% in 2002 vs. 48.9% in 2010, p<0.001) and specialists (35.2% in 2002 vs. 21.0% in 2010, p<0.05) also did not change significantly between 2005-2010 (p=0.55). Additionally, 80.12% of patient visits involving PPI prescriptions without an appropriate indication resulted in a 2.44-fold increase in PPI cost. The results of multivariable logistic regression revealed that, patients with more than 3 chronic conditions were significantly more likely to be prescribed a PPI without an appropriate indication as compared to patient visits with a chronic condition (Odds Ratio (OR) = 2.49, 95% Confidence Interval (CI): 1.26 - 4.94). CONCLUSIONS: PPI use among the elderly significantly increased over the study period, with 8 out of 10 patient visits involving PPI prescriptions without an appropriate indication. Additionally, the study found no change in PPI prescribing trends across physician specialties. Future research is required to identify the determinant variables associated with LCP use.

PHS105
IMPORTANCE OF JOINT WORK BETWEEN PHARMACIST AND PHYSICIAN IN THE RATIONAL DRUG USE IN PATIENTS WITH OSTEOPOROSIS
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OBJECTIVES: to estimate the tangible savings generated when there is a joint effort between Pharmacist and Physician METHODS: observational descriptive study in a tertiary care hospital patient records belonging to an 18 years old diagnosed with Osteoporosis in parenteral drug therapy with bisphosphonates (Zoledronic Acid 5mg and Botnic Acid 3mg). Observation period: June 2012 to June 2013 (n=161). The Pharmacist identified patients with non-pertinence to parenteral bisphosphonates. Non-pertinence was defined when: the patients had osteopenia (no additional risk factors), had renal function <35 ml/min, no previous scaling with oral medications and had previous osteoporosis. Finally, the pharmacist defined the behavior to follow (changing drug therapy). Analysis: were used absolute and relative frequencies, measures central tendency and dispersion. Treatment direct costs were quantified before and after assessment by the Pharmacist and The Physician. The statistical software SAS 21 license under the CES University was used. RESULTS: pharmacist non-pertinence identified in 34% (54/161) of patients, of which, the Physician agreed in 70% of cases (38/54). The 39% of patients switched to oral bisphosphonates Calcium Carbonate 70mg, Strontium Ranelate 2mg, Risedronate 35mg, 40% Calcium and Vitamin D3 and 21% left without medication. Given that patients were treated with Zoledronic Acid 5mg (25/38) or Botnic Acid 3mg (13/38), this conduct involved a decreased 18% a year later, with savings/ year $164 (considering only the value of the drugs). CONCLUSIONS: the joint work between Pharmacist and Physician showed an important saving in rational use of medicines.

PHS106
INCIDENCE OF POLYPHARMACY AMONG EMERGENCY PATIENTS AT A TERTIARY CARE HOSPITAL IN KARACHI: AN IGNORED PARADOX FOR QUALITY DRUG THERAPY
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OBJECTIVES: The aim of the study was to collect information of polypharmacy and its strength of association with respect to medication regimen among patients (pediatric and adults) visiting ED of a tertiary care hospital in Karachi Pakistan. METHODS: Retrospective study, conducted on all patients at Emergency Department (ED) of Aga Khan University Hospital, Karachi (AKUH) during the year 2012. The detailed clinical records on medication prescription from admission through discharge of all patients was reviewed. RESULTS: Total 51,000 patient ED visits during ED 2012 till December 2012, out of those polypharmacy was common in 40% of patients. Male were 56.6%(8,337) while 43.4%(7,553) were females. Pediatric patients were 17.9%(3,145) while 79%(14,279) were adults. The most common triage category for patients was F3 with 37.2%(6,483). Most of these patients were those who were recommended admission in other wards 59.6%(10,146), 26.5%(4,514) discharged patients and 9%(1,536) leave against medical advice (LAMA) patients. CONCLUSIONS: The perils and problems associated with Poly pharmacy are a subject of interest as polypharmacy was significant finding among all ED patients. The results from this study serves as a baseline to identifying the drug-related problems among ED and could helpful for pharmacists and physicians to develop and implement strategies to improve the quality of care. Keywords: Polypharmacy, Emergency Department, Emergency patients.

PHS107
PREVALENCE AND DETERMINANTS OF LOW-COST GENERIC DRUG PROGRAM USE IN THE PRIVATELY-INSURED ADULT POPULATION
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OBJECTIVES: Low-cost generic drug programs (LCPs) offer an affordable way for patients to obtain prescription drugs, but they may lead to exposure misclassification in administrative claims datasets. This study sought to assess the prevalence and determinants of LCP use in a privately-insured adult population. METHODS: This study relied on data from the Medical Expenditure Panel Survey (MEPS). The study was designed by two stages. (1) The total cost of the drug was paid out of pocket and 2) The cost of the drug exactly matched the cost of an LCP program. Demographics of LCP users and non-users were compared. A multivariable logistic regression model identified the determinant variables associated with LCP use. RESULTS: Of the total study...