May 2015 Work Group Recaps:

For results of Data Element Request Forms (DERFs) and External Code Set (ECLs) reviewed see DERF Resolution at http://www.ncpdp.org/members/members_wg_info.aspx?wgid=wgmc.

Work Group 1 Telecommunication

Ballot Adjudication:

- Ballot WG010065 for Telecommunication Standard (DERF 001224, 001213) is considered a Valid Ballot having received over 60% of Consensus Group votes. The Work Group adjudicated the comments received. Both comments were categorized as non-persuasive. Unless one of the commenters changes their vote, there will be a recirculation ballot. If the commenter changes their vote, the ballot will proceed to the Board of Trustees for approval after the required appeal period.

DERFs/ECLs Reviewed:

- DERF 001240 was pended.
- DERF 001274 was pended.
- DERF 001275/ECL000176 was approved.
- DERF 001276/ECL 000177 was approved.
- DERF 001277/ECL 000178 was approved.
- DERF 001278/ECL 000179 was pended to the WG1 FAQ Task Group.
- DERF 001279 was withdrawn.
- DERF 001280 was pended to a new MC Task Group.

Old Business:

- A HIPAA Standards, Operating Rules, and regulations (HPID/ICD/NPI/EFT/ERA) update was given.
- Use of Quantity Prescribed (46Ø-ET) in the Telecommunication Standard claim billing - Note this field would be required for Part D Schedule II Controlled Substance claims; however the use of this field is not limited to Part D claims only.
  - Telecom D.0 and all versions from that point have been updated (November 2012).
  - 03/2014 NCPDP received a response from HHS to the 03/2013 letter. The change will go through NPRM and Final Rule process.
    - We have asked the industry to have input on the implementation timeframe before the NPRM is published.
    - We have asked for a timeframe of NPRM publication.
  - 06/2014 Update: OESS has responded that an NPRM including Quantity Prescribed and SCRIPT electronic prior authorization is going through the review process.
  - WG1 Telecommunication FAQ Task Group brought forward a recommendation timeframe for Quantity Prescribed regulation implementation for OESS.
  - 04/17/2015 Update from NSG: At this time the Quantity Prescribed issue is going through the regulatory process. We will provide a revised target date for a regulation very soon.

- NCPDP SNIP Committee met twice in March and reviewed the analysis document and the timeline related to transitioning to an updated version of the HIPAA Telecommunication standard. The committee is in a holding pattern until the date for submission to the DSMO for consideration of the updated standard is determined. NCVHS is holding hearings in June on the 16th and 17th and have provided a list of questions they want NCPDP to respond to. SNIP will be creating a survey to obtain some of the responses they want NCPDP to respond to. SNIP will be reviewing and responding to questions. The committee will need assistance from WG45 and WG11.

Task Groups:
• The Telecommunication FAQ Task Group discussed questions submitted. There were no Editorial Guide requests this quarter. The group also made suggestions for some reorganization of the Telecommunication Implementation Guide.

• The Coordination of Benefits Task Group discussed questions submitted. Question 49 was reviewed, approved and it will be published in the Editorial Document. 12 COB related DERFs are being developed for submission in August.

• The Financial Information Reporting Task Group reviewed and updated FAQs, discussed the 36 month ATBT process including timing for quarterly retriggers, developed a retrigger timeline, updated the Post ATBT Whitepaper and updated the Plans Moving Processors Whitepaper.

• Information Reporting Problems Task Group received approval of the SPAP ADAP TrOOP Attestation document. They are creating a white paper to document best practices for matching N transactions, an SPAP/ADAPs Data Exchange White Paper, and a resource guide for Part D sponsors to help clean up overlapping OHI. They are also reviewing and updating the COB White Paper and processes when a B1 is reversed and the need to internally reverse the associated N1 prior to receiving a N2 request from the transaction facilitator.

• The Post Adjudication Task Group received a question on how to report 340B claims that were originally adjudicated as non-340B claims in the standard. A joint task group with the WG9 340B Task Group has been formed to address the question.

• The Definition of a Valid Prescriber Task Group brought forward Questions D36, O7 and O8. The task group developed a provider outreach letter to be used by stakeholders. The group has developed a separate FAQ document for questions related to the implementation of the 2015 Medicare D Final Rule for prescriber enrollment and improper prescribing practices. Questions are sent to CMS for their guidance.

• Part D Supplemental Payment Reporting Task Group continues to work on Nx report modifications. A reject code guide and a FAQ Document will also be developed.

• Eligibility Verification Enhancements Task Group determined the Last Known Segment is needed for the next HIPAA standard. The Task Group will go on hiatus until there new work items are brought forward or there are any new actions for Phase II of “Plan Types” for the message field or Last Known Segment.

• The Transaction ID Task Group is examining the use of a unique transaction identifier for all paid and approved Telecom transactions. They brought forward DERF 001274.

• Vaccine Services Task Group did not meet. This task group was disbanded.

• Benefit Integration Task Group did not meet.

• Standardized Subrogation Task Group reviewed Medicaid, Medicare Part D, and Commercial subrogation scenarios. A review of the request and response data elements for scenarios was completed.

Work Group 2 Product Identification

Ballot Adjudication:

• WG020005 for Billing Unit Standard is considered a Valid Ballot having received the required 60%+ of Consensus Group votes and 75%+ approval rating. No comments were received. After a 30-day appeal period, the ballot will be sent to the Board of Trustees in June 2015 and if approved will be published on the website in July 2015.

• WG020006 for Product Identification Standard is considered a Valid Ballot having received the required 60%+ of Consensus Group votes and 75%+ approval rating. No comments were received. After a 30-day appeal period, the ballot will be sent to the Board of Trustees in June 2015 and if approved will be published on the website in July 2015.
DERFs/ECLs Reviewed:

- DERF 1281 requests “Updated verbiage to the Product Identifiers Standard.” The DERF was approved.
- DERF 1282 requests “Updated verbiage for FAQ 7.42 of the Billing Unit Standard.” The DERF was approved.
- DERF 1283/ECL 180 requests “Updates to Appendix B within the External Code List (ECL).” The DERF/ECL was approved.
- DERF 1284 requests “Add FAQ to Billing Unit Standard to support the adjudication of QUIC form 201507 - Natesto (intranasal testosterone gel).” The DERF was approved.

Task Groups:

- The **Structure Product Labeling Activities Task Group** tracks the activities of the SPL, offers suggestions to improve access and usability of the FDA Structured Product Label and Electronic Drug Listings, and monitors the work of the Guiding Coalition for feedback to the work group. The Coalition has sent 19 letters to the FDA and is working on one additional letter of recommendation regarding storage conditions.
- The **Product Review and Billing Unit Exception Task Group** is reviewing the exceptions within the Implementation Guide and issues that result from changes to existing products or release of new products. The task group scripted and approved a letter to Golden State Medical Supply regarding GSMS’s inappropriate re-use of NDC Core 9 (Labeler Code + Product Code) for products having different active ingredients or strengths on at least three different occasions. WG2 approved this letter and it will be sent once it completes the NCPDP approval process. The task group also reviewed six new QUIC forms and submitted five of them to WG2 for final adjudication.
- The **NCPDP Product Identification Standard Task Group** developed the NCPDP Product Identifiers Standard Implementation Guide and have successfully balloted version 1.0 and 1.1 of that standard. DERF 1281 to add clarifying verbiage to the standard was submitted and approved at the May 2015 WG meeting. The Product Identifiers Standard Implementation Guide Version 1.2 will be balloted at the end of the August 2015 WG meetings.
- The **SPL REMS Requirements Task Group** is to gather the data needed to develop a template for pharmaceutical manufacturers to use in electronic submission of all the components for risk evaluation and mitigation strategies (REMS) drugs to a central repository (DailyMed) via FDA’s Structured Product Labeling system. The HL7 SPL Release 6 ballot that included support for REMS in the Common Product Model (CPM) was released last September. Comments received have been addressed and the ballot has passed. This greatly enhances FDA’s ability to move forward. Additionally, an Educational Track Session was held at this conference on the SPL initiative on REMs. Presenters were Ed Millikan of ASHP, Gerry McEvoy of ASHP, and Adam Kroetsch of FDA.
- The **Dates Associated with Pharmaceutical Products Task Group** investigates definition inconsistencies, involves government agencies to make them aware of the issues, and provides education on the importance via a white paper or other means. 55% of the white paper is completed but manufacturer input and participation is needed to finalize.
- The **Naming Standards for Drugs, Biologics and Biosimilars Task Group** sent a letter on February 27, 2015 to Sylvia M. Burwell, Secretary, HHS and Margaret A. Hamburg, MD, Commissioner, FDA on Nonproprietary Naming of Biologics, Biosimilars, and Interchangeable Biologics: Zarxio® (filgrastim) in which NCPDP expressed concerns about deviating from standard naming practices (e.g., USAN, INN) for biosimilars. They reported the FDA on Friday March 6, 2015, approved the drug Zarxio by Novartis' Sandoz unit as a lower-cost biosimilar alternative to Amgen's Neupogen. The FDA designated a placeholder nonproprietary name for this product as “filgrastim-sndz” but stated this designation of a nonproprietary name does not reflect on the agency’s yet-to-be-
published decision on a comprehensive naming policy for current and future biosimilar and other biological products. The task group scripted and approved a letter to United States Pharmacopeial Convention (USP) requesting their support and guidance for using recognized standard naming practices for biologics. The letter was approved by WG2 and it will be sent once it completes the NCPDP approval process. The task group also plans on scripting a letter to United States Adopted Names (USAN) Council and a follow-up letter to Secretary of Health with a copy to FDA Commissioner.

- The **Review of Appendix B Reference Code Qualifiers Task Group** is to review the definitions of existing product identifiers for accuracy and update as appropriate. The task group completed their review of the values and definitions within Appendix B and submitted DERF 001283 which was approved at this meeting. The ECL document will be updated and published after approval by the Board of Trustees.

- The **Application of the Billing Unit Standard Clarification Task Group** continues to make progress in identifying the rationale used to determine the billing unit from past QUIC forms/products reviews and is capturing the rationale/reasons for those decisions. Once the forms have been reviewed and the results compiled, a guidance document will be written.

- An update on the joint **WG11 and WG2 Drug Description Task Group** was not provided as they had not met.

- An update on the **MC NDC Depletion Task Group** was provided. See MC meeting minutes.

- An update on the **WG9 and WG2 Unbreakable Packages Task Group** was provided. See WG9 meeting minutes.

**New Business:**

**QUIC Form Review:**

- QUIC #201508 Signifor 20 mg NDC 00078-0641-81  
  **BU= EACH and Package Size=1** per sections 4.2.1 and 5.1.2 of the BUS
- QUIC #201509 Signifor 40 mg NDC 00078-0642-81  
  **BU= EACH and Package Size=1** per sections 4.2.1 and 5.1.2 of the BUS
- QUIC #201510 Signifor 60 mg NDC 00078-0643-81  
  **BU= EACH and Package Size=1** per sections 4.2.1 and 5.1.2 of the BUS
- QUIC #201511 Nascobal NDC 49884-0720-82  
  **BU=“Each” with a package quantity of 4 for the NDC 49884-0270-82 and a package quantity of 1 for the NDC 49884-0270-52 (inner pack)** per sections 4.2.1 and 5.1.12 of the Billing Unit Standard of the BUS
- QUIC #201512 ProAir Respliclick NDC 59310-0580-20  
  **BU=1 EA** based upon Section 5.4.1 of the BUS

**Work Group 3 Standard Identifiers**

**DERFs/ECLs Reviewed:**

DERF 001247/ECL 000175 – recommendation to MC to pend the DERF/ECL in order to review the request with WEDI’s Health ID Card Sub-Workgroup.

**Task Groups:**

- The **Pharmacy and/or Combination ID Card Task Group** presented initial draft language for the implementation guide regarding the use of smart phone technology for benefit identification. Work will continue during the next quarter to address discount card language, font size and guarantor vs. cardholder information. A DERF will be submitted for the August Work Group meeting requesting the implementation guide be revised to include this new section.

- The **Health Plan ID (HPID) Task Group** is on hold until CMS responds to NCVHS recommendations.
regarding the use of HPID in transactions.

Work Group 7 Manufacturer and Associated Trading Partner Transaction Standards

Old Business:
- WG7 received an update from WG9’s OIG Report OEI 05-12-00540 Task Group (see WG9 recap).

Task Groups:
- The Rebate Reference Guide Task Group did not meet this quarter.
- The Medical Rebate Standard Task Group did not meet this quarter.
- The Medicaid Drug Rebate Program Task Group identified the top three goals as ranked by the task group participants. Work began on Goal 1 which included an overview of the Medicaid Drug Rebates and Drug Rebate Equalization (DRE) Act. During the next quarter the task group will review the FAQ’s within the Implementation Guide and the Reference Guide to determine if those questions/answers should be combined in a standalone document.
- The Rebate Standard Update Task Group did not meet this quarter. During the next quarter the task group will focus on completing the vocabulary work, creating a workable solution for the integration of the Formulary and Benefit data and Specialty Pharmacy data exchange.
- The Regulatory Tracking/Pedigree Task Group did not meet this quarter.
- The Formulary Management Survey Task Group did not meet this quarter.

New Business:
- WG7 discussed the state of the current task groups and whether the needs of the Work Group are being met. The discussion will continue at the August Work Group meeting with the newly elected Co-Chairs.

Work Group 9 Government Programs

DERFs/ECLs Reviewed:
- 001285/Emergency ECL 000181 – recommendation to MC to approve the DERF/Emergency ECL.

Task Groups:
- The Prescription Monitoring Program (PMP) Task Group presented updated information for states that have prescription monitoring programs. The updated tracking document will be published.
- The 340B Task Group is working jointly with WG1 Post-Adjudication Task Group reviewing how “corrected” claims (designated as 340B) would be identified on a post adjudication file from an entity that accepts N1.
- The Medicare Part B Claim Billing for Dual Eligibles Task Group did not meet this quarter.
- The Health Insurance Exchange/Marketplace Task Group discussed CCIIO FAQ 10017 related to Special Enrollment Periods and retroactive effective dates and will explore using the subrogation process as the standard approach. The task group also reviewed a draft Commercial Coordination of Benefits Contractor proposal as it relates to the requirement for commercial carriers and Marketplace plans to coordinate benefits with ADAPs and accept payments for premiums and member copayments.
- The Medicaid Subrogation FAQ Task Group did not meet as no new questions were received.
- The Medicare Part D FAQ Task Group brought forward one question for review/approval by WG9 and two questions that were informational only. WG9 approved the response to the question. The FAQ document will be updated and published. The task group also requested WG9 review and approve the Post Adjudication PDE-DIR worksheet and guidance document. WG9 requested additional information be included in the guidance document. The task group will make those
revisions and present to WG9 in August. The task group is currently discussing reporting price concessions at point-of-sale and will have CMS participation in that discussion on a future call.

- **WG9/WG2 Unbreakable Package Joint Task Group** revised their objective, developed a spreadsheet to identify unbreakable packages by package type and setting of care (e.g. retail pharmacy, hospital, LTC, etc.) and worked on a draft letter to CMS which will recommend a NCPDP definition for unbreakable package.

- The **Supplemental Payer Part D Reconciliation Standardization Task Group** continued work on identifying data elements for a standard report, modified some field names due to anticipated changes for the next version of the Telecommunication Standard and worked on a draft user guide to accompany the standard report.

- The **Hospice Task Group** finalized a scenario grid based on guidance received and the multiple entities that need to communicate/process/respond to the guidance, submitted questions to CMS requesting clarification on hospice guidance, worked with the PDE sub-task group to identify a hospice scenario for post point-of-sale claim adjustment and worked with WG1 COB task group to define hospice for the Adjudicated Program Type field.

- The **Standardized Fraud, Waste and Abuse Training Attestation Task Group** completed review of all comments received on the standard attestation form. The task group continued work with NCPDP as they consider the role of gatekeeper for the standardized FWA attestation form as part of the existing database and the implementation timeframe should that occur.

- The **OIG Report OEI 05-12-00540 Task Group** finalized their scope and goals, developed a grid of options/potential solutions and reviewed adjudicated program types related to manufacturers.

**New Business:**

- WG9 received a Legislative update.

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**Work Group 10 Professional Pharmacy Services**

**DERFs/ECLs Reviewed:**

- DERF 001286 requesting changes to the Structured and Codified Sig Implementation Guide related to Dose Form was approved with no modifications.

**Task Group Reports:**

- The **MTM Communications Task Group** continues to develop new functionality using the CCDA release 2.

- The **Acetaminophen Best Practices Hospital Safety Sub-Task Group** had presented completed the content of the white paper that enumerates best practice guidelines, acetaminophen awareness, tools and technology solutions in a hospital model. During final clean-up additional supporting information was identified and is being incorporated. The paper will be brought back to WG10 for final approval. Suggestions were solicited for hospitals to participate in the planned study project.

- **WG11 Specialty Requirements for ePrescribing Task Group** is continuing the work to identify Specialty needs and brought forward DERF 001265 to support ePrescribing for Specialty drugs in the SCRIPT transactions.

**New Business:**

- Report on results of Ballot WG110064 adjudicated by WG11 that included the WG10 MTM changes to the Specialized Standard from DERF 001236 and the substance use DERF 001248. The ballot was approved and will go to the NCPDP Board of Trustees for approval after a 30-day appeal period. See WG11 for more information.
Donna Bohannon, R.Ph, CPPS Scientific Liaison, Nomenclature, Safety and Labeling Healthcare Quality Standards, United States Pharmacopeia: Current and planned projects of the USP including the Allergy Project.

**Work Group 11 ePrescribing & Related Transactions**

**Ballot Adjudication:**

- Ballot WG110064 for the SCRIPT and Specialized Standards is considered a valid ballot having received the required 68.21% of Consensus Group votes. Thirty-five comments were received. See Letter Ballot Comment spreadsheet for the ballot results. The ballot will go to the NCPDP Board of Trustees for approval after a 30-day appeal period.

- Ballot WG110065 for the Formulary and Benefit Standard is considered a valid ballot having received the required 68.21% of Consensus Group votes. Two negative with comments and one object with comments were received. See Letter Ballot Comment spreadsheet for the ballot results. If the submitters agree to change their comment to approve or abstain as appropriate, the ballot will not be recirculated and will proceed to the Board of Trustees for approval after the required appeal period. If the submitters do not wish to change their comment to approve or abstain as appropriate, the ballot will be recirculated.

**DERFs/ECLs Reviewed:**

- DERF 001237 was approved as modified by the Formulary and Benefit Task Group.
- DERF 001258 was denied.
- DERF 001259 was denied.
- DERF 001260 was denied.
- DERF 001261 was denied.
- DERF 001262 was pended for additional work by the Formulary and Benefit Task Group.
- DERF 001264 was approved as modified by the Electronic Prescribing Best Practices Task Group.
- DERF 001270 was approved as modified by the Specialty Task Group.
- DERF 001271 was pended for additional work by the Formulary and Benefit Task Group.
- DERF 001273 was approved.
- DERF 001280 was pended with the recommendation that a joint WG1/WG11 Task Group be created in MC Maintenance and Control.
- DERF 001287 was approved.
- DERF 001288 was pended.
- DERF 001289 was approved.
- DERF 001290 was pended.
- DERF 001291 was approved.
- DERF 001292 was approved.
- DERF 001293 was approved.
- DERF 001294/ECL 000182 was approved.
- DERF 001296 was approved.
- DERF 001297 was approved.
- DERF 001298 was approved.
- DERF 001299 was approved with modifications.
- DERF 001300 was pended.
- DERF 001301/ECL 000183 was approved with modifications.
- DERF 001302 was approved.

**Old Business:**
An industry update was provided on NCVHS Subcommittee on Standards, CMS (eprescribing), and DEA eprescribing for controlled substances and implementation activities.

Task Groups:
- The **Formulary and Benefit Task Group** brought forward DERFs 001237, 001258-001262, 001271, 001273, 001289, 001296-001302.
- **XML Task Group** reviewed submitted DERFs and brought forward recommendations.
- **NCPDP/HL7 Pharmacist Functional Profile Task Group** update was not given but report may be found in the May 2015 WG11 download file.
- **WG11 Electronic Prescribing Best Practices Task Group** provided recommendations for the SCRIPT Implementation Recommendations document on new terms for NCI Thesaurus, additional language for Diagnosis, zero refills approved on renewal and State Controlled Substance Registration Number. They also brought forward DERF 001264 and 001294/ECL 000182.
- No update was given from the **WG14 LTPAC ePrescribing Task Group**. Their task group recap may be found in the May 2015 WG11 download file.
- **REMS and ePrescribing Task Group** brought forward DERF 001288.
- The **Electronic Prior Authorization Workflow to Transactions Task Group** brought forward DERFs 001291 and 001292.
- **WG11/2 Joint Drug Description Task Group** no report was given.
- **The Meaningful Use and NIST Test Methods for ePrescribing Task Group** update was not given but the report may be found in the May 2015 WG11 download file.
- **Implementation of Structured Sig Task Group** provided recommendations for the SCRIPT Implementation Recommendations document on oral liquid medications. They also brought forward DERF 001287 and received approval to request a new NCI Thesaurus for DoseUnitOfMeasure.
- **Specialty Requirements for ePrescribing Task Group** requested input from WG participants to identify additional stakeholders. They asked participants to provide examples of information, workflow, etc. to inform discussion. They brought forward DERF 001265 to support ePrescribing for Specialty functions.

**WG14 Long Term and Post Acute Care (LTPAC)**

Old Business:
- Industry/Regulatory updates were provided which included HIPAA, NCVHS, and information from a recent Medicare Part D and LTC Listening Session

Task Group Reports:
- **The ePrescribing Task Group** – This Task Group are looking at electronic prescribing solution for IV administration.
- **The LTPAC Current Billing Issues Task Group** – The Task Group began looking at the NCPDP Telecommunication Standard to determine if the long term and post-acute care settings needs are met.
- Received updates from the **WG9 Hospice Task Group** and the **Medicare Part D FAQ Task Group**.

New Business:
- Heard a request from the **MC Education, Legislation and Regulations Task Group** soliciting help from LTC for comments being prepared for the ONC 2015 Certification Criteria NPRM.
- Heard information about a survey being created by the **NCPDP SNIP Committee** relating to HIPAA transactions. The survey results will be used to create testimony for the National Committee on Vital and Health Statistics’ (NCVHS).
Work Group 16 Property & Casualty/Workers Compensation

DERFs/ECLs Reviewed:
- DERF 001278 was reviewed with changes recommended to WG1.
- DERF 001295 was approved.

Old Business:
- An IAIABC update was provided.

Task Group Reports:
- The Legislative/Regulatory Monitoring and Education Task Group provided an update on state regulatory and legislative initiatives affecting Workers’ Compensation programs.
- The Billing and State Reporting Task Group provided an update regarding states moving to adopt regulations for e-billing, standard paper billing and EDI reporting. Presided over the DERF review.

New Business:
- Discussion on merging the two task groups into a single task group entitled Public Policy, Standards and Education Task Group. The discussion on task group realignment and responsibilities will continue at the next meeting in association with the annual review of work group scope and goals.
- New webinar task group was discussed but no action was taken.
- Perry Lewis of CoverMyMeds provided a brief talk on the value of NCPDP participation.

Work Group 45 External Standards Assessment, Harmonization and Implementation Guidance

Old Business:
- Industry updates were provided for WEDI, ASC X12, and CAQH CORE.

Task Groups:
- The Document Revision Task Group met once to review a change to the CARC mapping document for use of CARC 45 for paper claims and to update the NCPDP Procedure for updating the CARC Mapping Document to reflect changes in the Code Committee review process. The change to the mapping document was approved by the work group and will be go through the inter SDO process.
- The 834/835 FAQ Task Group received no new questions.
- A DSMO Task Group received no DSMO requests for review.
- CAQH CORE Task Group did not meet.

New Business:
- Issues surrounding the billing and reporting of the LA Provider fee were introduced and will be explored.

MC Maintenance and Control

Ballot Adjudication:
- Ballot WG010065 for Telecommunication Standard (DERF 001224, 001213) is considered a Valid Ballot having received over 60% of Consensus Group votes. The Work Group adjudicated the comments received. Both comments were categorized as non-persuasive. Unless one of the commenters changes their vote, there will be a recirculation ballot. If the commenter changes their vote, the ballot will proceed to the Board of Trustees for approval after the required appeal period.
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DERFs/ECLs Reviewed: (see DERF Resolution http://www.ncpdp.org/members/Work-Group.aspx?ID=wgmc)

• 41 new and pended DERFs/ECLs were reviewed (see WG1, WG2, WG3, WG10, WG11 and WG16 above).

• The DERFs approved at this meeting will result in 4 ballots for the August 2015 ballot period
  o WG020007 for Product Identifier Standard v12
  o WG100007 for the Structured and Codified Sig v21
  o WG110066 for SCRIPT
  o WG110066 for Formulary and Benefit v45

Old Business:

• Updates are available in the MC May 2015 download file:
  o HIPAA
  o NCPDP Legislative/Regulatory Activities

Task Groups:

• The Education/Legislation and Regulations Task Group will meet to provide comments on the NPRMs for Certification and Test Procedures

• The NDC Scarcity Task Group was disbanded.

• The PDMP White Paper Task Group reported on the activities of the S&I Framework PDMP pilots and received approval for their updated white paper.

To participate in the PDMP & Health IT Integration Initiative you can “join the initiative” by completing the form on the PDMP & Health IT Integration Initiative Join wiki page.

• The Unique Device Identifier (UDI) Task Group update was not given, but Task Group Report and presentation may be found in the May 2015 MC download file.

• The Real Time Benefit Check Task update was not given, but Task Group Report and presentation may be found in the May 2015 MC download file.

Project Development Form:

• Reviewed the pended New Project Development Form 000039 and approved a recommendation to create a Task Group to update the NCPDP Connectivity Standard.

New Business:

• The attendees received recaps of each Work Group’s activities.