May 2014 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/Work-Group.aspx?ID=wgmc

Work Group 1 Telecommunication

DERFs/ECLs Reviewed:

- DERF 001153/ECL 000148 was pended to WG1 Coordination of Benefits Task Group at the November and February Work Group. The DERF/ECL was recommended to MC as approved with modifications.
- DERF 001172/ECL 000152 was pended to WG1 Compound Billing Solutions Task Group at the February Work Group. The DERF/ECL recommended to be approved with modifications to MC.
- DERF 001188 was pended at the February Work Group for the WG1 Benefit Integration Task Group to continue work on the standard. The DERF was pended again for further work to be completed.
- DERF 001190 was approved.
- DERF 001191/ECL 000156 was pended for more work by the WG1 Vaccine Services Task Group.
- DERF 001192 was approved.
- DERF 001193/Emergency ECL 000157 was recommended to be approved to MC.
- DERF 001194/Emergency ECL 000158 was recommended to be approved to MC.
- DERF 001195 was approved.
- DERF 001196 was approved.

Old Business:

- The 2013 WG1 Year in Review was presented.
- 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 that the industry have input on the implementation timeframe before the NPRM is published.
  - We have asked for timeframe of NPRM publication.
  - Industry implementation was planned for January 1, 2014 – Notifications to industry, in Telecom D.0 Imp Guide, in Version D Editorial
    - This date is ON HOLD. OESS is working through options, but believes that a NPRM and final rule will have to be used.
- NCPDP SNIP Committee did not meet but the ICD-10 white paper has been updated with new information.

Task Groups:

- The Telecommunication FAQ Task Group discussed questions submitted. They discussed DERF 001196 (Hyphens). They reviewed DERF 001169 (ePA). They submitted a DERF for August on a modification to a value of Prescription Origin Code.
- The Coordination of Benefits Task Group discussed questions submitted. They will continue work on identification of benefit type for claims processing.
- The Financial Information Reporting Task Group reviewed reject code documentation and updated plan/processor recommendations, discussed industry recommended mass retrigger timing change approved by CMS. They reviewed a process for requesting next
plan of record for large volumes added to Post ATBT white paper, and brought forward FAQs for the FIR FAQ document.

- **Information Reporting Problems Task Group** is working on consolidating the data within the NCPDP SPAP ADAP BIN PCN list into a single worksheet. They are preparing a SPAP ADAP TrOOP Attestation document.
- The **Post Adjudication Task Group** brought forward DERF 001195 for a file layout for Transition Fill Claims Transfers.
- The **Audit Task Group** reviewed the NPRM for audit points to share with WG9. As their work is complete, they disbanded.
- The **Safe Use Processing – REMS Task Group** will remain active in case there are implementation questions or action items from WG11 ePrescribing and REMS Task Group.
- The **Definition of a Valid Prescriber Task Group** worked on responses to the 2015 Medicare D NPRM. They brought forward DERF 001193/Emergency ECL 000157 and 001194/Emergency ECL 000158. They are working through questions and have reached out to the DOJ for an understanding of the prescriber enrollment workflow. The task group is building a white paper of recommendations and gaps.
- The **Supplemental Payer Reporting Task Group** reviewed a Part D Plan recommended reporting for Nx transactions.
- The **Eligibility Verification Enhancements Task Group** worked on a project plan for a HIPAA demonstration project request. They are asking for input from industry to commitment to a pilot project or if the industry should just wait to implement the Last Known 4Rx Segment in the next round of HIPAA.
- The **Compound Billing Solutions Task Group** submitted DERF 001172/ECL 000152 for new Reject Codes and clarification to DUR/PPS Level of Effort. They submitted guidance for the Version D Editorial document.
- The **Transaction ID Task Group** is examining the use of a unique transaction identifier for all Telecom transactions. They are working through scenarios of best practices for Reversals. They seek input to the draft documentation.
- The **Vaccine Services Task Group** is completing recommendations for the Version D Editorial for pharmacy benefit billing for products, services, and products and services. They seek input to the documentation.
- The **Benefit Integration Task Group** is reviewing requirements for a standard for plans to share information on deductible, copay and Out of Pocket (OOP) to correctly maintain the Maximum out of pockets as described in the ACA. They continued updating pended DERF 001188 for industry awareness and participation.

**Work Group 2 Product Identification**

**DERFs/ECLs Reviewed:**
- DERF 001197 was approved with modifications.
- DERF 001198 was approved with modifications.

**Old Business:**
- The 2013 WG2 Year in Review was presented.

**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 18 letters to the FDA and is working on three additional letters of recommendation.
- 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.
  - Reviewed two new QUIC forms.
  - A Checklist of Impacts by Billing Unit Changes was completed.
One DERF requesting changes to the Billing Unit Standard was submitted for WG review.

- The SPL REMS Requirements Task Group is gathering 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 REMS Integration Initiative submitted REMS data elements and attributes to the FDA for review and submission to HL7. HL7 advised FDA on April 25, 2014 that the proposed REMS elements and attributes were out for balloting as part of the SPL ballot material. This is a major milestone in achieving the goal of standardization and codification of REMS via SPL.
- 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. The group continues to investigate dates used throughout the industry.
- The Evaluation of BUS’s Billing Units Task Group is gaining an understanding of the issues by looking at the unique attributes of patient administered products, identifying any challenges in the current BUS, reviewing billing and reimbursement processes for these products, identifying similarities amongst these products and reviewing past work on these products by WG2 and WG7. A presentation with speaker notes was completed and the task group was disbanded.
- The Naming Standards for Drugs, Biologics and Biosimilars Task Group met with FDA representatives on May 1, 2014. FDA suggested that they consider NCPDP developing the naming standard for biosimilars. Although this is not what NCPDP was seeking, it does give a good indication of the progress that was made during this meeting. There was discussion about an existing standards process for the naming of drugs in this country which is the USAN and the INN for international. WG2 proposed the FDA not deviate from this standardized process which may cause disruptions/adverse consequences in downstream processes.
- The Unique Device Identifier (UDI) Definitions Task Group completed their goal of defining the UDI as it applies to all applicable NCPDP standards and the information was passed to Maintenance and Control where a task group was formed to incorporate the UDI into existing NCPDP standards. The task group was disbanded.
- 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.
- An update on the joint WG11 and WG2 Drug Description Task Group was provided. See WG11 meeting minutes.
- An update on the MC NDC Depletion Task Group was provided. See MC meeting minutes.

New Business:

- Review of Appendix B Reference Code Qualifiers Task Group was formed to review and update as necessary the definitions of the Product/Service Qualifiers in Appendix B of the External Code List.
- QUIC Form Review:
  - #201406 Nicazel Doyx60 Kit NDC 42783-0610-60
    Per the standard, FAQ 7.6 and Section 4.2.1, the BU=each with a billing quantity of 120. The work group also discussed Nicazel Doyx30 NDC 42783-0610-30 and determined the BU=each with a billing quantity of 90 per the same references to the BUS.
  - #201407 Evzio NDC 60842-0030-01
Per Section 5.2.2 of the BUS, the package size should be 0.4 X 2 = .8 and the BU= mL and it is not considered an exception per FAQ 7.21. The trainer is the third unit and is excluded.

**Work Group 3 Standard Identifiers**

**DERF Reviewed:**
- DERF 001199 was approved.

**Old Business:**
- WG45 Provider Enrollment Task Group reported that ASC X12 is still working on their next version of the standard. It appears that the 274 will be used for provider enrollment and the 838 for EDI enrollment. During the past month the ASC X12 274 group began forming recommendations for the changes requested by NCPDP.
- The 2013 WG3 Year in Review was presented.

**Task Groups:**
- The **Pharmacy and/or Combination ID Card Task Group** finalized language for the implementation guide regarding the use of keychain cards as identification for the discount card program. The task group submitted DERF 001199 requesting this revision to the NCPDP Health Care Identification Card Pharmacy and/or Combination ID Card Implementation Guide.
- The **Pharmacy ID Card Operating Rules Task Group** did not meet this quarter.
- The **Health Plan Identifier (HPID) Task Group** – Ballot MC000006 passed and after a standard 30-day appeal period will go to the Board of Trustees for approval and release. The work of this task group is complete and the task group was disbanded.

**New Business:**
- Mapping Pharmacy ID Card to ASC X12 270 – WG3 approved the modifications to the mapping document. The document will be posted on WG3’s web page.
- Proposed Health Plan Certification Sub-Task Group – NCPDP is forming a task group under WG45 to review and draft comments on three Health Plan certification forms created by CAQH Core. It is expected that three to four conference calls will be necessary to review the forms and prepare comments. Health Plans were encouraged to participate.

**Work Group 7 Manufacturer and Associated Trading Partner Transaction Standards**

**Old Business:**
- The 2013 WG7 Year in Review was presented.

**Task Groups:**
- The **Reference Guide Task Group** did not meet this quarter.
- The **Medical/Biologics Task Group** did not meet this quarter.
- The **Medicare Part D Coverage Gap Task Group** was disbanded.
- The **CMS Task Group** did not meet this quarter. A new task group leader is being sought.
- The **Rebate Standard Update Task Group** is working on challenges identified with the implementation of version 07.00 of the Rebate Standard.
- The **Formulary Management Survey Task Group** did not meet this quarter.

**New Business:**
- A new sub-task group was formed under WG7’s Rebate Standard Task Group to standardize data sets exchange (utilization/clinical) between specialty pharmacy and manufacturers to support programs and agreements between the parties.
Work Group 9 Government Programs

Old Business:
- The 2013 WG9 Year in Review was presented.

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 formed a sub-task group to draft another educational letter to the State Medicaid Pharmacy Administrators.
- The Dual Eligible Recipients and Medicare Advantage Plans Task Group did not meet this quarter.
- The Health Insurance Exchange/Marketplace Task Group brought forward questions and recommended responses for review and approval by Work Group 9. The initial release of the FAQ document will be published.
- The Medicaid Subrogation FAQ Task Group did not meet as no new questions were received.
- The Medicare Part D FAQ Task Group brought forward questions and recommended responses for review and approval by Work Group 9. The FAQ document will be updated and published.
  - The Data Sharing for Overutilizers Sub Task Group. The white paper “NCPDP Recommendations for a Standardized Process to Share Medicare Part D Opioid Overutilization Data Between Sponsors” and the Standardized Overutilization Data Sharing template are being modified based on new CMS guidance.
  - The Supplemental Payer Part D Reconciliation Standardization Task Group continued to identify data elements for the reconciliation report and will work on developing a guide for the reconciliation report.
- The Hospice Task Group developed a form allowing communication between Hospice and Part D sponsors, “Hospice Status and Plan of Care for Medicare Part D A3 Reject Override.” The task group is developing instructions for use of the form.

Work Group 10 Professional Pharmacy Services

Old Business:
- The mL white paper “NCPDP Recommendations and Guidance for Standardizing the Dosing Designations on Prescription Container Labels of Oral Liquid Medications” was published and has been the subject of articles in the “Pink Sheet”, the American Pharmacists Association and an upcoming article in the Journal of the American Medical Association. In addition a two page call to action flier has been developed. The white paper (under Education/White Papers) and the flier (under Resources) are available to the public on the NCPDP web page at www.ncpdp.org.
- The MTM guidance document “Recommendations for Use of the HL7 Consolidated CDA Templates for Pharmacy” is now available on the NCPDP web page under Resources/Guidance Documents.
- The 2013 WG10 Year in Review was presented.

Task Group Reports:
- The MTM Communications Task Group continues with development of pharmacy targeted applications for existing Consolidated CDA documents. The CDA R2 MTM Templates R1, I2, US Realm passed the HL7 ballot with about 20 minor comments. The comments will be resolved and the standard will be moved forward for publication.
- The Acetaminophen Best Practices Task Group is continuing efforts to bring together the appropriate entities to develop a paper that enumerates best practice guidelines, acetaminophen awareness, tools and technology solutions in a hospital model.
- The work of the Specialty and Compounding Pharmacy Service Task Group has been divided into three groups: one in WG11 one in WG7 and one referred to WG1. The task group has not met and has no remaining items to address, therefore it was
The mL White Paper Task Group has completed the white paper, see Old Business above. The task group will remain open for another quarter in case further action is needed.

The Prescribable Medication Information at Point of Care Task Group has initiated a white paper aimed at recommending best practices to assure the availability of accurate and timely medication information at the point of care. The group has encountered difficulty getting supportive, non-anecdotal information and input from affected providers. It will be suspended pending discussion with the Co-Chairs regarding the best direction to take to address the effort.

Work Group 11 ePrescribing & Related Transactions

Ballots:
- Recirculation Ballot WG110058R for SCRIPT, Specialized (DERF 001123, 001146 and 001147) is considered a Valid Ballot having received the required 60%+ of Consensus Group votes and 75%+ approval rating. No new comments were received. See Letter Ballot Comment spreadsheet for the ballot results. The ballot will be sent to NCPDP Board of Trustees for approval after a 30-day appeal period.
- Ballot WG110059 for SCRIPT (DERF 001157, 001158, 001159) is considered a Valid Ballot having received 60% of Consensus Group votes. The Work Group adjudicated the comments received. One comment was voted as persuasive and editorial, so a modification will be made. The ballot will proceed to the Board of Trustees for approval after the required appeal period.

DERFs/ECLs Reviewed:
- DERF 001169 was pended to WG11 Prior Authorization Workflow to Transactions Task Group at the November and February Work Groups. The DERF was pended as there was not consensus on proceeding without further review. As this is the third pend, the DERF will not proceed. A new DERF may be submitted; the ePA Task Group will discuss whether the industry wishes to proceed with the work of this DERF with the addition of the pharmacy based workflow and bring forward a new DERF.
- DERF 001175 was pended at February Work Group for more review by the WG11 Formulary and Benefit Task Group. The DERF was approved with modifications.
- DERF 001176 was pended at February Work Group for more review by the WG11 Electronic Prescribing Best Practices Task Group. The DERF was approved with modifications.
- DERF 001178 was pended at February Work Group for more work by the MC Diabetes Task Group and the WG11 XML Task Group. The DERF was approved with modifications.
- DERF 001179 was pended at February Work Group for more work by the MC Diabetes Task Group and the WG11 XML Task Group. The DERF was approved with modifications.
- DERF 001180 was pended at February Work Group for more work by the MC Diabetes Task Group and the WG11 XML Task Group. The DERF was approved with modifications.
- DERF 001181 was pended at February Work Group for more work by the MC Diabetes Task Group and the WG11 XML Task Group. The DERF was approved with modifications.
- DERF 001182 was pended at February Work Group for more work by the MC Diabetes Task Group and the WG11 XML Task Group. The DERF was withdrawn.
- DERF 001183 was pended at February Work Group for more work by the MC Diabetes Task Group and the WG11 XML Task Group. The DERF was withdrawn.
- DERF 001184 was pended at February Work Group for more work by the MC Diabetes Task Group and the WG11 XML Task Group. The DERF was approved with modifications.
DERF 001185 was pended at February Work Group for more work by the MC Diabetes Task Group and the WG11 XML Task Group. The DERF was approved with modifications.

DERF 001200 was pended for more review by the XML Task Group.

DERF 001201 was pended for more discussion as part of the real-time benefit check.

DERF 001202 was approved with modifications.

DERF 001203 was approved.

DERF 001204 was approved with modifications.

DERF 001205/ECL 000159 was recommended to MC for approval.

DERF 001206 was approved.

DERF 001207 was pended for more discussion of the Formulary and Benefit Task Group.

DERF 001208 was approved with modifications.

Old Business:

- An industry update was provided on NCVHS Subcommittee on Standards, CMS (eprescribing), and DEA eprescribing for controlled substances and implementation activities.
- The 2013 WG11 Year in Review was presented.

Task Groups:

- The Formulary and Benefit Task Group submitted DERF 001207, 001208 and worked on pended DERF 001175.
- XML Task Group reviewed submitted DERFs. They reviewed the MC Diabetes Task Group DERFs and brought forward recommendations.
- NCPDP/H77 Pharmacist Functional Profile Task Group did not meet this quarter.
- WG11 Electronic Prescribing Best Practices Task Group worked on pended DERF 001176 (WrittenDate/EffectiveDate), submitted DERF 001202 and provided recommendations for the SCRIPT Implementation Recommendations document on the use of WrittenDate/EffectiveDate and on CancelRx processing.
- An update was given from the WG14 LTPAC ePrescribing Task Group. The group focused its efforts on capturing industry conventions for use of SCRIPT 10.6 in long-term and post-acute care settings, for use by facility vendors and pharmacies that are migrating HL7 interfaces to the SCRIPT Standard (to meet the Medicare ePrescribing standards).
- REMS and ePrescribing Task Group has refined their use cases between prescriber, pharmacy, REMS Administrator, switch/intermediary for safe use programs, and the data elements needed. They are working through flows from basic REMS to complex REMS exchanges. They will work through the actual transactions and then updates to an implementation guide. They have provided trigger/data element analysis information to the WG2 SPL Task Group.
- The RxFill Task Group brought forward updates to the SCRIPT Implementation Recommendations document for the Fill Status Notification transactions. As their work is completed, they disbanded.
- The Electronic Prior Authorization Workflow to Transactions Task Group met to discuss pended DERF 001169 and new DERF 001201.
- The WG11/2 Joint Drug Description Task Group continued work on a project with NLM for creating processes for supporting RxNorm eprescribing names. They are working through programmable rules for injectable names. They brought forward additional guidance for the SCRIPT Implementation Recommendations document.
- The Pharmacy to Pharmacy Prescription Transfer Task Group was monitoring Ballot WG110058. As their work is completed, they disbanded.
- The Meaningful Use and NIST Test Methods for ePrescribing Task Group is discussing testing for RefillRequest/Response, RxChange and other transactions.
- **Implementation of Structured Sig Task Group** began building guidance on implementation and Q&A from implementers. They have collected top used sigs, created examples of those sigs, and are building guidance.

- **Specialty Requirements for ePrescribing Task Group** is discussing the electronic prescribing flow and data needs and what might already be supported in the SCRIPT transactions.

- **Prescription Delivery Task Group** has completed their work and disbanded

**New Business:**

- The attendees began discussion on beginning the process to name the next version of SCRIPT for electronic prescribing since the regulatory process and the implementation timeline take a while.

- Information was provided that Minnesota is reconvening their eRx work group to update the eprescribing guidance that was published prior to the 01/01/2011 mandate. The group will be reviewing the guide to reflect more current state, as well as providing information about some of the newer transactions that aren’t yet widely adopted (Cancel, Change, etc.)

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

**Old Business:**

- Industry/Regulatory updates were provided which included HIPAA and NCVHS.

- The 2013 WG10 Year in Review was presented.

**Task Group Reports:**

- The **ePrescribing Task Group** reported they are preparing guidance on the use of SCRIPT transactions in LTC. The LTPAC HIT Summit will be held June 22-24, 2014 in Baltimore, MD. The Minnesota eHealth Advisory Committee has approved an ad-hoc eRx workgroup to update existing ePrescribing Guidance.

- The **LTPAC Current Billing Issues Task Group** discussed a pharmacy workflow which included the use of the ePA transactions. They prepared comments to the included in the NPRM for 2015 Medicare Part D changes affecting the LTPAC settings.

- Received updates from the **WG1 Eligibility Verification Enhancements Task Group**, the **WG9 Medicare Part D FAQ Task Group** and the **WG9 Hospice Task Group**.

**Work Group 16 Property & Casualty/Workers Compensation**

**Old Business:**

- An IAIABC update was provided.

- The 2013 WG16 Year in Review was presented.

**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. The task group has completed development of content for a WEDI white paper to explain the handling of the special requirements for workers compensation in pharmacy billing.

**Work Group 45 External Standards Assessment, Harmonization and Implementation Guidance**

**Old Business:**

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

- The 2013 WG45 Year in Review was presented.

**Task Groups:**

- The **Document Revision Task Group** presented examples of vaccine billing and ASC X12 reporting. The examples were approved contingent on approval by WG1 as part of the Version D Editorial document. Modifications to the CARC mapping document that had
been presented at the February work group meeting were approved for publication. The ASC X12 Request for Pharmacy Industry Survey involving provider initiated payments to payers has been finalized and will be distributed shortly. The task group reviewed examples developed by the WG1 Compound Billing Issues Task Group and found no needed changes to the WG45 documents. Based on a request from the task group for direction, the work group approved a motion for the task group to begin looking at the CARC/RARC mapping at a detailed level rather than a consolidated level as is currently done.

- The 834/835 FAQ Task Group received no new questions, therefore did not meet.
- A DSMO Task Group received no new DSMO Change Requests. The task group met to review the ASC X12 changes to support the Health Plan Identifier (HPID) and provided input to the WG3 public comment response.
- The Provider Enrollment Task Group did not meet as information was received that the move from the ASC X12 274 to the ASC X12 838 Standard applied only to EDI enrollment. The previously submitted ASC X12 Change Request is actively being reviewed at this time.
- The Central Pay Task Group completed its work prior to the last work group meeting. The document was reformatted from the spreadsheet to a standard guidance document and is moving through the Inter SDO approval process. Once approved the task group will disband.

MC Maintenance and Control

Ballots:
- Recirculation Ballot WG110058R for SCRIPT, Specialized (DERF 001123, 001146 and 001147) is considered a Valid Ballot having received the required 60%+ of Consensus Group votes and 75%+ approval rating. No new comments were received. See Letter Ballot Comment spreadsheet for the ballot results. The ballot will be sent to NCPDP Board of Trustees for approval after a 30-day appeal period.
- Ballot WG110059 for SCRIPT (DERF 001157, 001158, 001159) is considered a Valid Ballot having received 60% of Consensus Group votes. The Work Group adjudicated the comments received. One comment was voted as persuasive and editorial, so a modification will be made. The ballot will proceed to the Board of Trustees for approval after the required appeal period.

DERFs/ECLs
34 new and pended DERFs/ECLs were reviewed (see WG1, WG2, WG3 and WG11 above). The DERFs approved at this meeting will result in:
- Five ballots for the May 2014 ballot period
  - WG010062 for Telecommunication vE6
  - WG010063 for Post Adjudication v44
  - WG02004 for Product Identification Standard Implementation Guide v10
  - WG110060 for SCRIPT, XML and Specialized
  - WG110061 for Formulary and Benefit v42

Old Business:
- Updates given:
  - HIPAA
  - NCPDP Legislative/Regulatory Activities
- The 2013 MC Year in Review was presented.

Task Groups:
- The Education/Legislation and Regulations Task Group prepared comments for meaningful use regulation.
- The Ordering of Diabetic Supplies Standard Task Group worked with the WG11 XML Task Group to resolve issues with the DERFs submitted at the February WG. Two of the DERFs were withdrawn because the functionality exists today and the other DERFs passed and will go to ballot.
The NDC Depletion Task Group held a preliminary call with FDA Office of Compliance staff to begin discussion on concerns and suggestions for future solutions. The task group will begin working on a white paper of best practices for use of product identifiers and how to address changes after the approval of the Product Identification Standard.

The PDMP White Paper Task Group reported on the activities of the S&I Framework PDMP group. Currently the S&I Framework PDMP stakeholders are reviewing standards and their possible gaps. Three standards are standing out. They include the ASAP web service, the PMIX PDMP and the SCRIPT Medication History. The next step in the process after the gaps have been resolved is creating an implementation guide which is where participation from the PDMP White Paper Task Group and WG11 will be important. 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.

New Business:
- The attendees received daily Work Group recaps.
- New Project Development Form #000038 requesting the creation of a new task group to look at creating a standard attestation form was approved for recommendation in WG9. The Standardization Co-Chairs will review and present determination at the August WG.