AAPS Open Forum Report: Proposals for Regulatory Harmonization of a Global BCS Framework: Challenges and Opportunities

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An open forum entitled Proposals for Regulatory Harmonization of a Global BCS Framework: Challenges and Opportunities was held at the AAPS Annual Meeting and Exposition in San Diego, CA, on November 4, 2014.

The first speaker was Dr. Mehul Mehta (Division of Clinical Pharmacology I, FDA) whose presentation was entitled The Criteria for BCS-Based Biowaiver and the Regulatory Experience Gathered in the US. Dr. Mehta began by outlining the basis for BCS-based biowaivers, related criteria, implementation, and outcomes. He referred to the guidance released by FDA in August 2000 (1). The criteria include the requirement that initial in vivo bioavailability characterization is needed for NDAs and that BCS consideration is not applicable for these studies. Whereas BCS consideration is applicable for waivers of bioequivalence (BE) studies in NDAs (pre- and post-approval) and ANDAs, the drug substance has to be BCS Class 1, the product has to be rapidly dissolving, plus the test and reference formulations should be pharmaceutical equivalents and show rapid and similar dissolution. The highest strength (HS) should be soluble in 250 mL or less of aqueous medium over the pH range of 1–7.5, it should be highly permeable (HP) where it should have a 90% or greater bioavailability or urinary recovery or permeability greater than the reference compound(s), and it should be rapidly dissolving (RD) where 85% or more dissolves in 30 min at pH 1, 4.5, and 6.8. Additional requirements include consideration of excipients used, prodrugs, and narrow therapeutic index (NTI) drugs; drug products that are absorbed from the oral cavity are excluded.

Dr. Mehta then presented a decision tree where the primary consideration is whether the sponsor asked for a BCS 1 classification; if not, the reviewer decides if implementation can continue. If acceptable, implementation involves submission of the summary package to the committee by the primary reviewer for review where each member has one vote. The choice is “YES,” “NO,” or “insufficient information for BCS 1 classification,” and the decision is by majority vote. An official record of each consult consisting of the summary report, discussion, vote, and outcome is kept. The decision is then communicated back to the review team and back to the sponsor via the review division.

Dr. Mehta informed the audience that the committee has met several times a year to review submitted applications. Sixty-three drug products came up for evaluation, 42/63 (67%) were classified as BCS Class 1, 30/63 were from the New Drugs side, 17/30 (57%) received Class 1 determination, and 33/63 were from the OGD side where 26/33 (79%) were classified as BCS Class 1. On the New Drugs side, 15/30 were at the IND Stage 8 and received Class 1 determination and agreement on biowaivers, five were turned down, and two had insufficient information. Fifteen (15/30) were at the NDA review Stage 9 and received Class 1 determination and related regulatory relief, two were turned down, and four had insufficient information. On the Generic Drugs side, 26/33 cases received BCS Class 1 determination, five were turned down, and two had insufficient information. Fifty seven out of 71 ANDAs were subsequently approved.

Dr. Mehta then informed the participants that guidance updates were being considered, as follows:

• Biowaivers for BCS Class 3 drugs with Q1, Q2 restrictions.
• Addition of “very rapid” dissolution criteria (>85% in 15 min)
• Change permeability boundary from 90% to 85%.
• Change the pH solubility range from 1–7.5 to 1–6.8.
• Change “highest dosage strength” to “highest strength.”
• Dissolution medium volume of 500 mL instead of 900 mL; possibility of changing paddle speed from 50 to 75 rpm.
• Additional topics and clarification on FDCs (fixed dose combinations), ODTs (orally disintegrating tablets), MR (modified-release) products.
• Update the list of model drugs.
• Strengthen GI stability requirement.

He indicated that the revised draft should be available for public comments in the next few weeks and summarized as follows:

• BCS guidance is founded on sound scientific and regulatory bases.
• CDER has created a centralized BCS evaluation process to assure consistency and transparency across all therapeutic areas and generic drugs.
• Since the issuance of the guidance in 2000, FDA has reviewed 63 drug products across NDAs and ANDA, of which 42 (67%) were given BCS Class 1 determination.
indicating wide application of the BCS principles in
drug development and evaluation; in terms of number
of applications approved, this would be more than 100
applications.
• In vivo studies are used and relied upon extensively for
permeability classification of drug substances.
• In vitro permeability studies can provide pivotal
information. They are often very helpful to resolve
uncertainties from in vivo data.
• Appropriate reporting of all necessary information (e.g.,
GI stability, method suitability) will help timely evaluation
of BCS classification submissions.
• The BCS guidance is being revised to allow biowaivers
for BCS Class 3 drugs as well; additional changes are
also being made to bring it into greater international
harmonization.
• The revised draft guidance should become available
within next few weeks (December 2014).

Dr. Mehta acknowledged contributions from Ramana
Uppor, Jaya Vaidyanathan, and Duong (Diane) Nhu.

The next speaker, Professor J-M. Cardot (University
of Auvergne), presented a talk entitled The Criteria for
BCS-Based Biowaiver and the Regulatory Experience
Gathered in the EU. The various conditions for a biowaiver
were described as follows:
• Highly soluble drug substances.
• Known human absorption.
• No narrow therapeutic index drugs.
• Immediate-release, solid oral dosage forms intended for
systemic action and having the same pharmaceutical
form.
Drug solubility using the highest therapeutic dose should
be tested over the pH range of 1.0–6.8 using a volume
of 250 mL, and a solubility–physiologic profile should be
constructed. The pH must be measured before and after
sampling (replicate points) using the shake-flask or other
justified method. Initial drug permeability considerations
are based on the fraction absorbed ($F_a$) and not based
on bioavailability ($F$). Candidates for biowaiver in the EU
should be completely absorbed as indicated by $F_a \geq 85\%$.
Reported bioequivalence between aqueous and solid
formulations via the oral route may be supportive since it
indicates that absorption limitations due to formulation
characteristics may be considered negligible, and if $F >
85\%$, no problems are expected. Additional permeability
considerations involve a mass balance study (provided no
gastrointestinal metabolism or degradation) where $>>85\%$
is absorbed. Metabolism, before or after permeation (i.e.,
reduction vs oxidation), and the mechanism of absorption
must be critically assessed. Well-performed in vitro
permeability investigations including reference standards
may also be considered as supportive of in vivo data.

Biowaivers are not applicable for sublingual, buccal,
and modified-release formulations; however, biowaivers
for orodispersible formulations are applicable when
absorption from the oral cavity can be excluded. Excipients
that might affect bioavailability must be qualitatively and
quantitatively the same, whereas other excipients must be
qualitatively the same and quantitatively very similar.

Professor Cardot then turned to the dissolution
requirements for a biowaiver. Here the sampling time
points must be performed at least every 15 min, and more
frequent sampling during the period of greatest change
in the dissolution profile is recommended. If dissolution is $>85\%$
in 15 min, no statistical calculations are needed, whereas
if dissolution is $>85\%$ but $<30$ min (BCS 1), $f_2$ calculations
or alternative tests are required. He emphasized that
generally, the risks of an inappropriate biowaiver should be
more critically reviewed (e.g., site-specific absorption, risk
for transport protein interactions at the absorption site,
excipient composition, and therapeutic risks) for products
containing BCS 3 as opposed to BCS 1 drug substances.

For fixed combination (FC) drugs, BCS-based
biowaivers are applicable for immediate-release FC
products if all the actives in the FC belong to BCS 1 or BCS
3 and provided that the excipients fulfill the requirements.
Otherwise, in vivo BE is required.

The European public assessment report EPARS 5
from June 2014 discussed the BCS-based biowaivers.
There are requirements for specific individual biowaivers
for miglustat and oseltamivir. The applicant must provide
evidence of the possibility of BCS-biowaivers for a product,
otherwise BE studies are required.

Professor Cardot ended his presentation with statistics
relating to biowaiver acceptances in the various European
countries, as follows:
• In Portugal, less than 1% of the submissions were based
on biowaivers.
• In Spain, 18 submissions were made for biowaivers and
15 were approved (14 were classified as BCS Class 1
and one as BCS Class 3) with three rejections. Of these,
11 were national applications (dossier submitted only
in one country and not in a European framework), two
centralized applications where another member state
was the assessor (European authorization route resulting
in a centrally authorized product with a single marketing
authorization), and two DCPs (European authorization
route resulting in a mutual recognition procedure, MRP)
from Germany. Professor Cardot explained that the
difference between MRP and DCP is that a product must
be previously authorized in at least one Member State on
a national basis for MRP to be used. DCP may be used if
the product has not previously been authorized in any
Member State, but the Sponsor does not want to use the
centralized procedure, or the product is not eligible for
the centralized procedure.
• In the Czech Republic, 16 biowaiver applications were
made, 11 were approved, and five rejected. There were
two national applications where one was approved and
one rejected.

- In Austria, only three products were submitted based on BCS Class 1 or 3 biowaivers.

Professor Cardot explained further that in some instances, the dossier may be reviewed and assessed by a Reference Member State (RMS) and then followed up by a Concerned Member State (CMS). One of the proposed Member States may be asked by the applicant to act as RMS where the RMS does the initial evaluation and then issues a draft assessment report. The other Member States (CMS) either agrees with the RMS evaluation or may ask further questions or raise objections.

The next presentation entitled Differences in Biowaiver Criteria that Can Pose Difficulties and Challenges for Global Registration of Generic Drug Products was made by Dr. Gerald Beuerle (Teva Ratiopharm, Ulm, Germany).

The various BCS guidances in the United States, Europe, and Canada were compared. The general requirements are follows:

- Applicable to immediate-release products where the drug substance is highly soluble.
- For orally administered drugs excluding narrow therapeutic index drugs and intended for systemic action; the test and reference may contain different salts.
- Solid drug products with the same pharmaceutical form and no absorption from the oral cavity.
- Absorption/permeability properties considered as well as excipient considerations with at least rapid dissolution.

The biowaiver criteria for products containing BCS Class 1 drugs include high solubility, complete absorption, contain the same excipients as the reference product, and dissolution >85% in 30 min. If the excipients might affect bioavailability (e.g., mannitol, surfactants), they should be qualitatively and quantitatively the same (in Canada, should be within ±10% of amount in reference product).

For products containing BCS Class 3 drugs, the drug must be highly soluble with limited absorption, contain the same excipients as the reference product, and have dissolution >85% in 15 min. If the excipients might affect bioavailability (e.g., mannitol, surfactants), they should be qualitatively and quantitatively the same, whereas if other excipients are used, then they should be qualitatively the same and quantitatively very similar.

Dr. Beuerle then described the solubility requirements in the European, Canadian, and United States guidances. He indicated that there were many similarities between the European and Canadian requirements compared with those of the United States. Firstly, in Europe and Canada, the highest dose should completely dissolve in 250 mL of relevant dissolution media, whereas in the United States, the highest strength should completely dissolve in 250 mL of relevant media. In Europe and in Canada, the pH range is 1–6.8 whereas in the United States, it is 1–7.5. Furthermore, in the former countries, dissolution in pH 1.2, 4.5, 6.8 (plus at pK_a for EU) should be tested whereas in the United States, dissolution testing is based on ionization characteristics (e.g., pH 1, 7.5, pK_a, pK_a+1, pK_a−1 if 3 < pK_a < 5). The same temperature (37 ± 1 °C) and shake-flask method as well as verification of pH before and after addition of drug substance are common requirements in all the jurisdictions. However, in Canada and Europe, replicate determinations may be necessary (in Canada, not less than three), whereas at least three replicate determinations in each pH are required in the United States.

Permeability/absorption requirements differ somewhat between Canada/Europe and the United States. In the former countries, complete absorption in humans is required, whereas in the United States, permeability can be determined in humans or with in vivo–in vitro intestinal permeability methods. The absolute BA absorption criterion in Europe and Canada is at least 85%, whereas in the United States it is at least 90%. Human data can be collected in mass balance or from absolute BA studies in all the jurisdictions; in Europe and Canada in vitro permeability is considered as supportive only, but in the United States in vitro data or an alternative to human studies are acceptable. For mass balance studies, stability under GI conditions needs to be demonstrated in Europe and Canada and published literature data may be acceptable, but in the United States, for some permeation studies, stability in the GI tract needs to be demonstrated and no mention is made of use of published data.

As far as the in vitro dissolution requirements are concerned, for BCS 1, at least 85% should be dissolved under physiologically relevant conditions and similar in vitro dissolution using f_2 calculations are required in all the jurisdictions. Also, there is no f_2 requirement if more than 85% has dissolved within 15 min. In Europe other suitable tests than f_2 are permitted but not mentioned in the Canadian and U.S. requirements. The investigations should be carried out at pH 1.2, 4.5, and 6.8 in all the jurisdictions and potentially in the pH of lowest solubility in Europe and Canada. In all the jurisdictions, SGF and SIF are permitted as alternatives, but surfactants are not permitted. Enzymes are permitted throughout only if the product contains gelatin (coatings/capsules), and paddle and basket speeds of 50 rpm and 100 rpm, respectively, are specified using a temperature of 37 ± 1 °C. A volume of 900 mL or less is specified for Europe and Canada, but only 900 mL in the United States. The sampling schedule of 10, 15, 20, 30, and 45 min are specified in Europe, and 5, 10, 15, 20, and 30 min in Canada. In the United States, the required sampling schedule is 10, 15, 20, and 30 min, and at least 12 units must be tested in all jurisdictions. In Europe, testing of more than a single batch is advisable, a minimum of two batches to be tested in Canada comprising at least a 100K/commercial size batch and documentation with the study protocol, and full validation must be provided in Europe and Canada. However, these two requirements are not mentioned in the U.S. regulations.
The use of certain excipients is described for BCS 1 drugs in Europe and Canada, where well-established excipients must be used with a description of their respective functions and should be the same excipients as in the reference product. In the United States, it is specified that for BCS 1 drugs, the excipients that are currently used in FDA-approved in IR solid oral dosage forms are acceptable. Furthermore, for BCS 1 drugs in Europe and Canada, the usual amounts of excipients with justification are specified, whether the amounts are in the normal range and are preferably similar as in the reference product. In the United States, the quantity of excipients should be consistent with the intended function. Any excipients that might affect bioavailability, such as sugar alcohols and surfactants, should be qualitatively and quantitatively the same in Europe, and must be within ±10% of the amount in the reference product in Canada. Large quantities of certain excipients, such as surfactants and sweeteners, may be problematic in the United States.

Dr. Beuerle contended that many requirements of the 2000 FDA guidance are still valid and have been included in the European and Canadian guidelines and that there appears to be even more advanced requirements in the latter countries guidelines. Several challenges for global registration of a BCS-based submission are suitability of BCS Class 3 drug products, different assessment of permeability, stricter criteria for certain excipients in EU/CAN, some other minor differences requiring additional efforts, and acceptance of the reference product used.

Dr. Jack Cook (Pfizer, United States) made the following presentation entitled Differences in Biowaiver Criteria that Can Pose Difficulties and Challenges for Global Registration of New Products. Dr. Cook began with a table (2) showing a provisional BCS classification of the top 200 drugs in the United States, United Kingdom, Spain, and Japan where 61% of those can be considered as potential biowaiver candidates (34 BCS 1 and 27 BCS 3). However, not all regulatory agencies will allow BCS-based biowaivers for these drugs. He referred to the differences in the definition of highly soluble between the FDA and the EMA. In the former instance, highly soluble is defined as when the highest dose strength is soluble in 250 mL or less of aqueous media over the pH range of 1–7.5 (at 37 ± 1 °C), but in the EMA it is the highest single dose strength administered as immediate-release formulation(s) and is completely soluble in 250 mL of buffers within the pH range 1–6.8 at 37 ± 1 °C. This difference in the definition of highly soluble reduces the potential biowaiver candidates to 34% of the tabled 200 drugs. Furthermore, the difference in the definition of highly permeable (HP) between the FDA and the EMA further reduces the potential biowaiver candidates to 27%. In the United States, the permeability class boundary is based indirectly on the extent of absorption, measurement of the rate across human intestinal membranes. Alternatively, non-human systems can be used, and the drug is considered HP when the extent of absorption is 90% or more, whereas the EMA considers a drug as HP when it is completely absorbed (i.e., where the measured extent of absorption is ≥85%), and complete drug absorption must be justified based on reliable investigations in humans. Dr. Cook estimated from a survey of 84–167 studies per annum, that using dissolution instead of conducting biostudies with typically 32 subjects per study, could result in a cost difference of approximately $220,000 per study. Because of differences in BCS biowaiver requirements among the various regulatory jurisdictions, he estimated that such differences lead to an approximate 70% reduction in BCS-based waivers (relative to maximal use), resulting in an increase in development costs and time AND unnecessary exposure and risk to human subjects.

Dr. John Gordon, a WHO representative, presented the next talk entitled BCS-Based Biowaivers: Who Criteria and Experience. He explained the purpose of the WHO guideline, which is based on one of the WHO constitutional functions to provide objective and reliable information and advice in the field of human health, a responsibility that it fulfills in part through its program of publications. The organization seeks, through its publications, to support national health strategies and address the most pressing public health concerns of populations around the world. He emphasized, however, that the WHO, unlike the FDA or EMA, is not a regulatory body. The WHO initiated the United Nations Prequalification Programme for Priority Essential Medicines with an action plan in 2001 for expanding access to selected priority medicines with the objective to ensure quality (Q), safety (S), and efficacy (E) of medicines procured using international funds such as GFATM, UNITAID, and so forth. This involves the evaluation of Q, S, and E of prioritized essential medicines, inspections of manufacturers, and monitoring of the products after their prequalification. It also includes prequalification of quality control laboratories and building capacity of regulators and quality control laboratories. The key output of the program includes a list of prequalified medicinal products for the treatment of HIV/AIDS, malaria, tuberculosis, influenza, neglected tropical diseases, acute diarrhea, and reproductive health. The program invites applications for a specified set of medicinal products identified on the program’s Expressions of Interest (EOI) and contained in the list of WHO Essential Medicines (EML) and identified by WHO treatment programs. The core EML list presents the minimum medicine needs for a basic healthcare system, listing the most efficacious, safe, and cost-effective medicines for priority conditions. Priority conditions are selected on the basis of current and estimated future public health relevance, and the potential for safe and cost-effective treatment. The complementary list presents essential medicines for priority diseases, for which specialized diagnostic or monitoring facilities, specialist medical care, and specialist training are needed. In case of doubt, medicines may also be listed as complementary on
the basis of consistent higher costs or less attractive cost-effectiveness in a variety of settings.

The WHO has published guidelines on registration requirements for multisource (generic) pharmaceutical products, Annex 7 (3), a proposal to waive in vivo bioequivalence requirements for WHO Model List of Essential Medicines immediate-release, solid oral dosage forms, Annex 8 (4), and also prequalification programme guidelines with general notes on Biopharmaceutics Classification System (BCS)-based biowaiver applications (2012) and BCS-based biowaiver applications for Reproductive Health (RH) products (2013). Annex 7 is currently under revision. An International Generic Drug Regulator’s Pilot Project (IGDRP) has been created to promote regulatory collaboration and convergence in generic drug regulatory programs to address challenges posed by increasingly heavy workloads, globalization, and the growing complexity of scientific issues. Participating countries were surveyed on their biowaiver practices.

Annexure 7 describes the criteria for API classification where high solubility is defined as follows:

- The highest single therapeutic dose is soluble in 250 mL or less of aqueous media over the pH range of 1.2–6.8.
- The pH–solubility profile of the API should be determined at 37 ± 1 °C in aqueous media.
- A minimum of three replicate determinations of solubility at each pH condition is recommended.

Dr. Gordon presented a table wherein countries including Australia, Brazil, Canada, European Union, Mexico, New Zealand, Singapore, South Africa, and Switzerland and WHO use the highest single therapeutic dose as a solubility criterion, whereas the United States and South Korea use the highest strength on the market.

The high permeability criteria for API classification include:

- Extent of absorption in humans is 85% or more based on a mass balance determination or in comparison with an intravenous dose.
- An acceptable alternative test method for permeability determination of the API could be in vivo intestinal perfusion in humans.
- When used for permeation studies, suitability of the methodology should be demonstrated, including determination of permeability relative to that of a reference compound whose fraction of dose absorbed has been documented to be at least 85%, as well as use of a negative control.

Supportive data can be provided by (1) in vivo or in situ intestinal perfusion using animal models, (2) in vitro permeation across a monolayer of cultured epithelial cells (e.g., Caco–2) using a method validated using APIs with known permeability, although data from neither method would be considered acceptable on a stand-alone basis. High permeability is assessed with respect to the permeability of a series of reference compounds with documented permeabilities and fraction absorbed values, including some for which fraction of dose absorbed is at least 85%.

A table was presented wherein countries including Australia, Brazil, Canada, European Union, New Zealand, Singapore, and Switzerland and WHO require human absolute bioavailability/mass pivotal and other supportive studies with ≥85%, and the United States and South Korea also accept in vivo intestinal perfusion and in vitro permeation studies but ≥90%.

Initially, the eligibility of an API for a biowaiver includes BCS Class 1, 2 (weak acids), and 3, but Annex 7 has recently been revised where only BCS Class 1 and 3 APIs are eligible. Also, excluded from biowaivers are narrow therapeutic index drugs.

A table showing that in Brazil, Singapore, South Korea, Taiwan, and the United States, only BCS Class 1 APIs were eligible, whereas in Australia, Canada, European Union, Mexico, New Zealand, South Africa, and Switzerland and WHO considered also BCS Class 3 APIs. In Japan, biowaivers based on the BCS was not accepted. Dr. Gordon provided examples of drugs and their respective BCS classifications for HIV/AIDS and related diseases (abacavir sulfate, Class 3; emtricitabine, Class 1; lamivudine, Class 3; stavudine, Class 1; and zidovudine, Class 1) and anti-tuberculosis medicines (ethambutol, Class 3; isoniazid, Class 3; levofloxacin, Class 1; moxifloxacin HCl, Class 1; ofloxacin, Class 1; and pyrazinamide, Class 3). Diethylcarbamazapine, used in neglected tropical diseases, is considered a BCS Class 3.

Conventionally, immediate-release products only are eligible and comparative dissolution profiles (CDP) are required. The following dissolution test conditions have been specified:

- Comparative testing should ensure the similarity of the test and comparator product in three different pH media considered relevant for absorption from the GI tract.
- Comparative in vitro dissolution testing should be conducted in at least three aqueous media of pH 1.2, 4.5, and 6.8.
- Volume of medium: 900 mL
- Temperature of medium: 37 ± 1 °C
- Agitation: paddle apparatus at 75 rpm or basket apparatus at 100 rpm
- Replicates: 12 units
- Sampling schedule: e.g., 5, 10, 15, 20, 30, and 45 min
- Surfactants not permitted–Revised (clarified)
- “Very rapidly” is defined as not less than 85% of the labeled amount is released within 15 min or less from the test and comparator product, and profile comparison is not needed.
- “Rapidly dissolving” means not less than 85% of the labeled amount is released within 30 min or less from the test and comparator product, and profile comparison (e.g., f2 testing) is required.

For BCS Class 1, the following are also considered:
**Excipients**

- Should employ well-known excipients in usual amounts
- Beneficial to contain similar amounts of the same excipients
- Critical excipients (e.g., mannitol, sorbitol, surfactants), if present, should not differ qualitatively or quantitatively

**Comparative in vitro dissolution**

- Products should be similarly rapidly dissolving
- NLT 85% in 30 min for both products
- $f_2$ profile comparison (unless 85% in 15 min for both FPPs)

For BCS Class 3, the following apply:

- APIs are highly soluble but limitations to absorption
- Excipients
  - Qualitatively the same excipients
  - Quantitatively very similar (as per Level 1 change according to SUPAC)
- Comparative in vitro dissolution
  - NLT 85% dissolved within 15 min for both products

Dr. Gordon then compared the agitation speeds used in different countries. For Apparatus 1 (basket) in Brazil, Canada, Mexico, Singapore, United States, and WHO, 100 rpm is specified whereas in Australia, the European Union, New Zealand, South Africa, South Korea, and Switzerland, usually 100 rpm is specified, and in Taiwan it is 50–100 rpm. When Apparatus 2 (paddle) is used, 50 rpm applies in Brazil, Canada, Mexico, Singapore, and United States, usually 50 rpm in Australia, the European Union, New Zealand, South Korea, and Switzerland, 50–75 rpm in South Africa and Taiwan, and 75 rpm by WHO.

There were various other considerations for biowaivers such as:

- BCS-based biowaivers for some FDCs difficult—comparator not available.
- FDCs must include only Class 1 or 3 APIs to be eligible (e.g., rifampicin containing products are not eligible for a BCS-based biowaiver)—only mono-component products eligible (e.g., United States and Mexico).
- Waiver for only one component of a fixed drug combination product (FDC) possible (e.g., Canada).
- Orodispersible tablets are eligible if there is no sublingual or buccal absorption (e.g., WHO, European Union, Australia, Brazil, and Canada, but Canada excludes products taken without water).

Since 2008, BCS-based biowaivers have been accepted by WHO for medicines for HIV/AIDS and related diseases, both monocomponent and FDCs, and for anti-tuberculosis medicines where levofloxacin and moxifloxacin have been the most frequent biowaivers.

Regarding pediatric strengths where there are no equivalent strength comparator products, the WHO, in consultation with experts in the area, is exploring the scientific basis for accepting biowaivers for pediatric products whose adult strengths are eligible for a BCS-based biowaiver. The principles involved for the API would be that a biowaiver would only applicable for eligible APIs on the list and include an additional solubility criterion to account for the smaller fluid volume in pediatric stomach (e.g., 50 mL). Furthermore, excipient comparisons should be conducted on a proportional scale, and comparative dissolution studies in media of at least three pH levels, both single and multiple unit comparisons.

The next presentation, entitled **BCS-Based Permeability: Managing Global Expectations**, was delivered by Dr. Gordon L Amidon (University of Michigan, Ann Arbor, MI). Dr Amidon emphasized that the labeling of a product is important since the product must do what the label states and the patient does not have a choice and must simply comply. The question he posed is, “How do we ensure labeling?” The ultimate reference for use of a product is the data resulting from the Innovator Phase 3 Product study. However, today, a multi-source product requires BE to be confirmed with the innovator commercial product. Dr. Amidon referred to the first labeling law in 1516 in Germany, which was related to the purity of German beer. As far as harmonization is concerned, he noted that that “world harmonization” would involve 196 countries. Small APIs are the same everywhere, and their chemistry well established. Their manufacture and shipping are well established, and the excipients used are usually well known, but are they all the same? He referred to some limitations with harmonization such as country-by-country sovereignty. Also, pharmacodynamics (efficacy) differences among different populations (genetic/epigenetic) with dosing implications, differences in absorption, distribution, metabolism and elimination (ADME), and cultural differences with implications for the route of administration, formulation (taste), and so forth. Hence, looking forward, scientific consensus is required, such as consideration by WHO and FIP, country sovereignty and related scientific training, and consequently BE harmonization and the need for in vivo predictive dissolution. He contended that the first steps should include consideration of oral immediate-release products, the establishment of worldwide scientific consensus within WHO/FIP and local societies, and country-by-country education.

Dr. Amidon then discussed the science of bioequivalence and stated that “The science of BE is at the absorption site.” He then described the movement of drug through the gastrointestinal tract and emphasized that pharmacopeias such as the *USP/EP/JP/ChP* must be more active in developing BE standards for oral products and the use of dissolution. There should be national cooperation and discussion/debate with the goal of human health. He referred to the FDA BCS guidance published in 2000 and the associated recommendations as well as the subsequent WHO Technical Report on the Investigation.
of Bioequivalence and the EMA Note for Guidance on the Investigation of Bioavailability and Bioequivalence.

Dr. Amidon explained that there were differences between pharmacopoeia and state (regulatory agencies such as FDA/EMA) initiatives and priorities with the inception of the USP in 1820 and the FDA in 1906. In 1938, the FDA Act was concerned with safety, and only in 1962 did amendments address efficacy. In 1984, the Drug Price Competition and Patent Term Restoration Act was published, which enabled the era of generic medicines, and in 1992, PDUFA was introduced to facilitate NDAs. He emphasized that the pharmacopoeias have tread lightly on the issue of BE and dissolution standards and that USP “compliance with any of the [dissolution] tests does not assure bioavailability or bioequivalence.”

Dr. Bhoopathy stated that the issue of BCS biowaivers is simple yet intricate, as the scientific framework is based on well-defined characteristics such as solubility, permeability, and dissolution. However, it is intricate since there is a lack of harmonization, and differences exist in interpretation and application. He referred to the “permeability divide” where Group 1 considers the classification based on solubility and permeability and requires human PK or intestinal permeability methods, whereas Group 2 bases classification on solubility and absorption plus human PK or reliable literature data.

Considering Group 1, high BCS (human in vivo) permeability is considered equal to high absorption (i.e., fraction absorbed or $F_{abs}$), and high in vitro $P_{app}$ is always equal to high BCS permeability, therefore high in vitro $P_{app}$ is always equal to high $F_{abs}$. However, high BCS permeability is not always equal to high in vitro $P_{app}$ due to barrier differences, transporter expression, experimental conditions, and so forth. Group 2 requires that complete absorption is demonstrated from human PK studies or published literature, and in vitro data are only considered as supportive. Bioavailability is not equal to $F_{abs}$ since many BCS 1 drugs are extensively metabolized. Furthermore, mass balance can be variable and inaccurate since <30% of BCS 1 drugs are highly variable, and there are many steps between formulation performance and site of measurement and limited subject numbers with only mean values reported. The quality of submissions may result in subjective interpretation of published data. Dr. Bhoopathy then raised the question why not accept permeability as pivotal. The reason may be that there could be insufficient reproducibility and also the impact of pH on evaluation plus a conflict of in vitro with known in vivo data. He then presented data to illustrate each of the foregoing reasons. In particular, there could be a misclassification by local permeability. Using the drug pindolol as an example, he showed that high permeability at any point along relevant intestinal regions can result in high absorption, and that a pH of 7.5 is the average pH of the human ileum, which accounts for about half of the small intestine length. He then cited examples of three compounds, where urine recovery was 57–79% and fecal recovery 15–21% for compound 1, urine recovery of 29–63% and fecal recovery of 18–38% for compound 2, and an absorption range of 74–100% based on urine recovery for compound 3. For all these compounds, the FDA recommends the biowaiver alternative implying that in vitro testing provides a more direct and accurate classification. He outlined some harmonization steps where for Group 1, highly permeable is when absorption is >85% and biowaivers for BCS 3 drug substances should be allowed. For Group 2, the use of permeability as definitive data should be allowed and begin with those drugs that are transported by passive mechanisms.
He concluded by showing a table of Global Acceptance:

<table>
<thead>
<tr>
<th>Country</th>
<th>Year Issued</th>
<th>Dissolution</th>
<th>Permeability</th>
<th>Solubility</th>
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<tbody>
<tr>
<td>United States</td>
<td>2000</td>
<td>85% in 30 min</td>
<td>90%</td>
<td>1–7.5</td>
</tr>
<tr>
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<td>2001 (2010)</td>
<td>85% in 15 min</td>
<td>85%</td>
<td>1–6.8</td>
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<tr>
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<td>2004</td>
<td>Rapid</td>
<td>High</td>
<td>1–6.8</td>
</tr>
<tr>
<td>WHO</td>
<td>2006</td>
<td>85% in 30 min</td>
<td>85%</td>
<td>1.2–6.8</td>
</tr>
<tr>
<td>ANVISA</td>
<td>2011</td>
<td>Specified List</td>
<td>85%</td>
<td>Specified List</td>
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<tr>
<td>TGA</td>
<td>2011</td>
<td>85% in 15 min</td>
<td>85%</td>
<td>1–6.8</td>
</tr>
<tr>
<td>Canada</td>
<td>2012</td>
<td>85% in 30 min</td>
<td>85%</td>
<td>1.2–6.8</td>
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REFERENCES

