Design Concept for a Confirmatory Basket Trial

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• Co-authors on the present work:

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- Pathway design subgroup, additional members:
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- Small populations workstream is part of **DIA Adaptive Design Scientific Working Group (ADSWG),** a group of 180 statisticians and clinicians from industry, academia, and national health authorities (FDA and EMEA)

Small Populations Within A Common Disease

- The increasing discovery of molecular subtypes of cancer leads to small subgroups that actually correspond to orphan or "niche" indications, even within larger tumor types
- Enrolling enough patients for confirmatory trials in these indications may be challenging.
- The shift to a molecular view of cancer requires a corresponding paradigm shift in drug development approaches
- Exclusive use of "one indication at a time" approaches will not be sustainable

Approaches to development based on predictive biomarkers

- Optimized co-development of a single drug and its companion diagnostic
 - Gives a clear hypothesis and answer and still has a role in selected instances
 - Will be challenging to do in niche indications
- "Umbrella" trials
 - One tumor type with multiple drugs and predictive biomarkers
 - Patients are matched to drugs based on predictive biomarkers
 - Cooperation among multiple sponsors
 - Examples: BATTLE, I-SPY, Lung-MAP
- "Basket" or "bucket" trials
 - Multiple tumor types with one drug and predictive biomarker
 - Approval based on pooled analysis
 - Premise is that molecular subtype is more fundamental than histology
 - Single sponsor

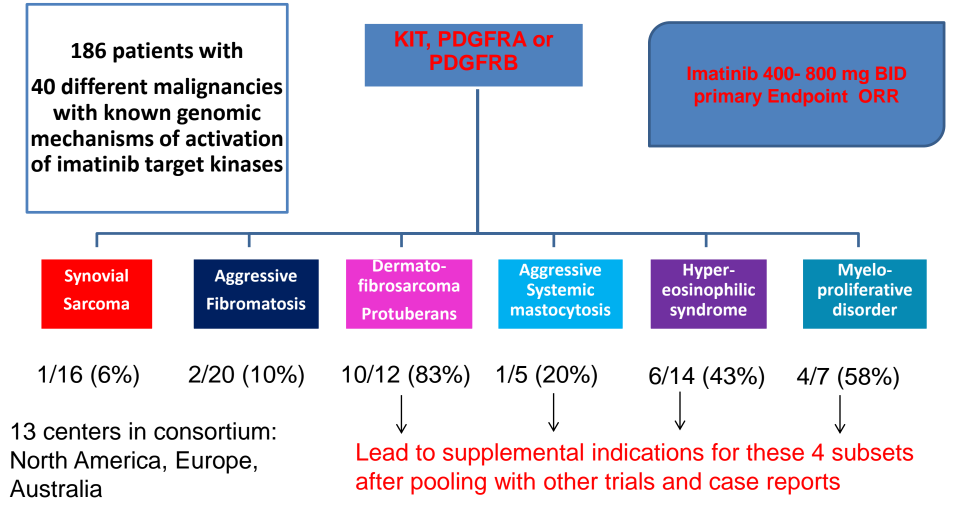
Agenda

Introduction

- General Design Concept for a Basket Trial
- Challenges of Basket Trials and Recommendations for Overcoming Them
- Detailed Design Considerations

Conclusions

The Original Basket: Imatinib B2225



April 2819 menthal. Innovative trial designs to accelerate the availability of highly effective anti-cancer therapies: an FDA perspective, AACR 2014

Basket Trials to Date

- A similar design to Imatinib B2225 was endorsed at a Brookings/Friends Conference in 2011
- Common features:
 - Exploratory and opportunistic in nature
 - Single-arm trials with ORR as primary endpoint
 - Intend to use pooled population for primary analysis to gain broader indication across tumor types (individual tumor type is not adequately powered)
 - Involve <u>possibly</u> transformative medicines in patients with great unmet need and <u>seemingly</u> exceptionally strong scientific rationale

Issues

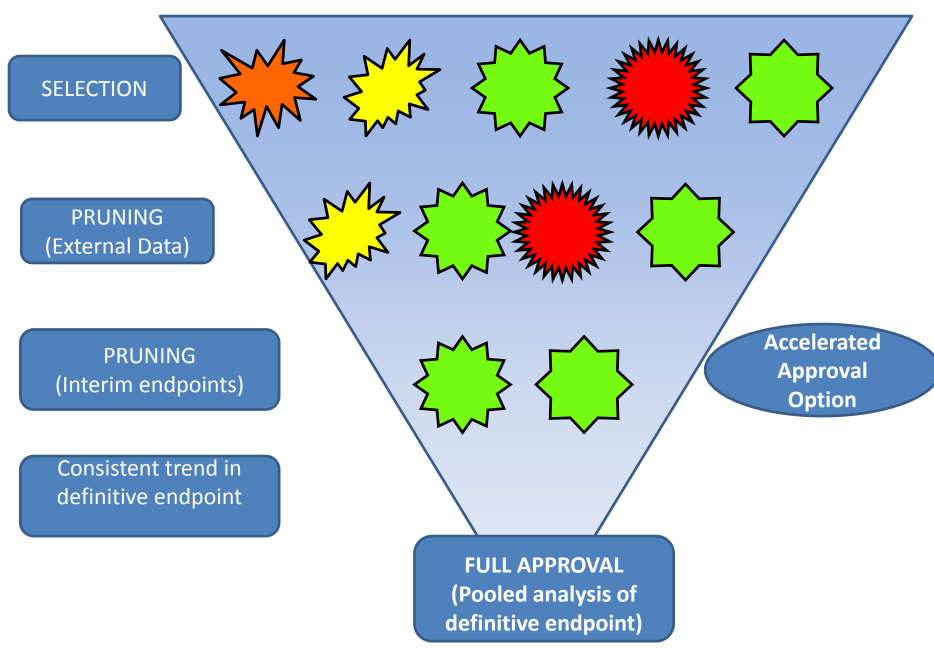
- Clinical data to support pooling my be limited, and treatment effect may differ between tumor types
 - Vemurafenib works in melanoma with BRAF V600E mutation but not colorectal cancer with same mutation
- Not all drugs hoped to be transformational live up to this promise
- Response rate may not predict overall survival
- Single arm trials are subject to patient selection bias
- Predictive effect of a biomarker is confounded with the prognostic value which is often unknown
- Health authorities can be non-committal upfront

DIA Small Population Pathway Subteam

- Can we develop a generalizable confirmatory basket design concept with statistical rigor?
 - Applicable not only to exceptional cases, but to all effective medicines in any line of therapy
 - Follow existing accelerated and standard approval pathways to increase drug approvability
- This would have multiple benefits
 - Increase and accelerate access to effective medicines for patients in niche indications
 - Provide sponsors with cost-effective options for development in niche indications
 - Provide health authorities with more robust packages for evaluation of benefit and risk

GENERAL DESIGN CONCEPT

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Features of the Design (I)

- Tumor histologies are grouped together, each with their own control group (shared control group if common SOC)
- Randomized control is preferred
 - Single arm cohorts with registry controls may be permitted in exceptional circumstances as illustrated by imatinib B225 and others
- In an example of particular interest, each indication cohort is sized for accelerated approval based on a surrogate endpoint such as progression free survival (PFS)
 - This may typically be 25-30% of the size of a Phase 3 study
- Initial indications are carefully selected as one bad indication can spoil the entire pooled result

Features of the Design (II)

Indications are further "pruned" if unlikely to succeed, based on:

- External data (maturing definitive endpoint from Phase 2; other data from class)
- Internal data on surrogate endpoint
- Sample size of remaining indications may be adjusted based on pruning
- Type I error threshold will be adjusted to control type I error (false positive rate) in the face of pruning
 - Pruning based on **external** data does not incur a statistical penalty
 - Discussed in more detail later in talk
- Study is positive if pooled analysis of remaining indications is positive for the primary definitive endpoint
 - Remaining indications are eligible for full approval in the event of a positive study
 - Some of the remaining indications may not be approved if they do not show a trend for positive risk benefit as judged by definitive endpoint

CHALLENGES OF BASKET DESIGNS AND RECOMMENDATIONS FOR OVERCOMING THEM

Challenge 1: Risks of Pooling

- One of more bad indications can lead to a failed study for all indications in a basket
- Histology can affect the validity of a molecular predictive hypothesis, in ways which cannot always be predicted in advance
 - Vemurafenib is effective for BRAF 600E mutant melanoma, but not for analogous colorectal cancer (CRC) tumors
 - This was not predicted in advance but subsequently feedback loops leading to resistance were characterized

Addressing challenge 1

- Basket trials are recommended primarily after there has been a lead indication approved (by optimized conventional methods) which has validated the drug, the predictive biomarker hypothesis, and the companion diagnostic
 - Example, melanoma was lead indication preceding
 Brookings trial proposal in V600E mutant tumors
- Indications should be carefully selected
- Indications should be pruned in several steps before pooling

Challenge 2: Adjusting for Pruning

- Pruning indications that are doing poorly on surrogate endpoints may be seen as cherry picking
 - This can inflate the false positive rate, an effect termed "random high bias"
- Addressing the challenge:
 - Emphasize use of external data, especially maturing Phase 2 studies, for pruning
 - Pruning with external data does not incur a penalty for random high bias
 - Apply statistical penalty for control of type I error when applying pruning using internal data
 - Methods for calculating the penalty are described in stat methods papers (see key references)
 - Rules for applying penalty must be prospective
 - Penalty is not large enough to offset advantages of design

Challenge 3: Will the companion diagnostic assay generalize across indications?

- Analytical properties of assay may depend on tissue type
- Cutoff between biomarker positive and negative may vary between tissue types for a continuous biomarker
- Addressing the challenge:
 - Analytical validation of the assay for all relevant indications prior to study start
 - Prior to study start, recommend biomarker stratified randomized phase 2 studies to set provisional cutoffs for continuous biomarkers in each indication to the extent feasible

Challenge 4: Availability of tissue

- Tissue sampling and processing are variables that can greatly affect the outcome of a study based on a predictive biomarker
- Basket studies will require cooperation and uniformity across departments organized by histology
- Addressing the challenge:
 - The sponsor must have extensive contact with the pathology department and relevant clinical departments at all investigative sites and provide standard methods for tissue sampling, handling, and processing
 - The sponsor should engage an expert pathologist who is dedicated to training prior to trial start, and troubleshooting during the trial

Challenge 5: Clinical validity of the predictive biomarker hypothesis

- The clinical validity of the predictive biomarker can only be verified by inclusion of "biomarker negative" patients in the confirmatory study
- Addressing the challenge
 - Recommend a smaller pooled, stratified cohort for biomarker negative patients, powered on surrogate endpoint
 - Would need to expand the biomarker negative cohort (to evaluate definitive endpoint) if surrogate endpoint shows possible benefit
 - Prior evidence should permit this if:
 - An approved lead indication has already provided clinical evidence for the predictive biomarker hypothesis
 - Prior phase 2 studies support the predictive biomarker hypothesis in other indications

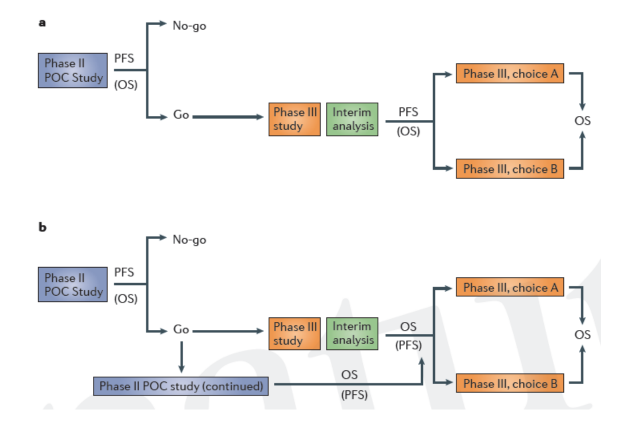
Challenge 6: High Screen Failure Rate

- Pro: patients will have access to tailored therapy
- Con: patient has a high risk of being a screen failure if biomarker positive subgroup is low prevalence
- Addressing the challenge:
 - Study should provide a broad-based test like NGS which will give the patient some guidance on alternative therapies if they are screen failures for basket study

Challenge 7: Interim endpoints may not predict definitive endpoints

- Addressing the challenge:
 - Prefilter indications based on maturing definitve endpoint data from phase 2
 - See Figure 2
 - Require consistent trend in definitive endpoint for final full approval

Phase 2 Influencing Phase 3 Adaptation: The Phase 2+ Method



Beckman, R.A., Clark, J. & Chen, C. Integrating predictive biomarkers and classifiers into oncology clinical development programmes. *Nature Reviews Drug Discovery* **10**, 735-748 (2011)

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Another Possible Source of External Data

- Real World Data (RWD) from Off-Label Use
- Impact of RWD on basket trial performance is currently under study in a project led by postdoctoral fellow Daphne Guinn



DETAILED DESIGN CONSIDERATIONS

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Designs to Be Compared

- Sample size changes after pruning
 - D0: No pruning and no change (benchmark)
 - D1: No increase to sample size after pruning
 - D2: Sample size in pooled analysis after pruning remains same as planned for the trial (SS)
 - D3: Sample size for trial remains same after pruning as planned for the trial (SS)

Designs	Overall Trial	Pooled Population				
DO	SS	SS				
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D2	>SS	SS				
D3	SS	<ss< td=""></ss<>				

Type I error control

- k tumor indications each with sample size of N and all with 1:1 randomization
- An interim analysis is conducted at information fraction *t* for each tumor indication and a tumor will not be included in the pooled analysis if p-value> α_t
- The pooled analysis will be conducted at α* so that the overall Type I error is controlled at α when there is no treatment effect for any tumor (H0)
- What is α^* ?

Solving for adjusted alpha (α^*)

- Let Y_{i1} be the test statistics based on information fraction t, and Y_{i2} be the test statistics based on the final analysis of data in the *i*-th cohort (i=1, 2,...,k)
- Suppose that *m* cohorts are included in the final analysis (*m*≥1), and let V_m be the corresponding test statistics. The probability of a positive outcome in pooled analysis is

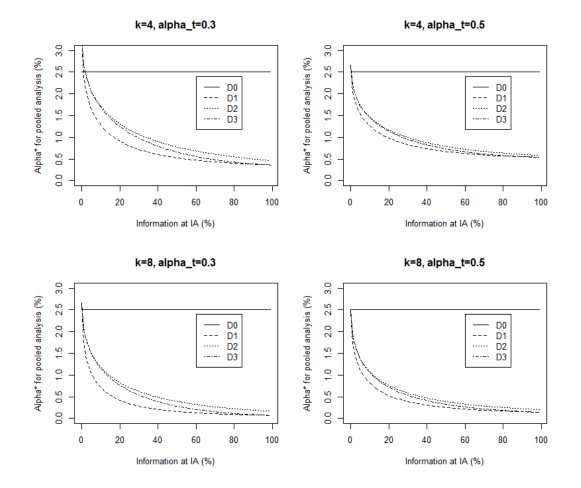
 $Q_0(\alpha^*|\alpha_t, m) = \Pr_{H_0}(\bigcap\{Y_{i1} > Z_{1-\alpha_t} \text{ for } i=1,...,m\}, \bigcap\{Y_{j1} < Z_{1-\alpha_t} \text{ for } j=m+1,...k\}, V_m > Z_{1-\alpha^*})$

or $Q_0(\alpha^*|\alpha_t, m) = \Pr_{H_0} (\cap \{Y_{i1} > Z_{1-\alpha_t} \text{ for } i=1,...,m\}, V_m > Z_{1-\alpha^*})(1-\alpha_t)^{(k-m)}$

• α * is solved from below where c(k, m) = k!/((k-m)!m!)

$$\sum_{m=1}^{k} c(k,m) Q_0(\alpha^* | \alpha_t, m) = \alpha$$

α^* under different design options



 α^* decreases with increasing k as expected, but its relationship with α_t is complicated with April 2017 the interplay between cherry-picking and futility stopping.

Comparison of operating characteristics

- k=6 tumor indications with total planned event size (kN) ranging from 150-350
 - The true treatment effect is –log(0.6), or hazard ratio of 0.6 in a time-to-event trial
- Pruning occurs at when half of the events have occurred
- Number of active indications (g) with target effect size ranges from 3 to 6, with remaining ones inactive

Study power and sample sizes under different pruning and pooling strategies

Planned	Number of	Power (%) for a			Exp. number of			Exp. number of			
events	active	positive study			events for pooled			events for overall			
	tumors				population			study			
		D0	D1	D2	D3	D0/D2	D1	D3	D0/D3	D1	D2
200	6	95	85	95	93	200	157	179	200	179	221
200	5	85	75	91	86	200	144	172	200	172	228
200	4	67	62	82	76	200	131	166	200	166	234
200	3	44	45	68	61	200	119	159	200	159	240
300	6	99	96	99	99	300	254	277	300	277	323
300	5	96	81	98	96	300	232	266	300	266	334
300	4	84	81	94	91	300	209	255	300	255	345
300	3	60	64	84	79	300	187	244	300	244	356

An Application of Special Interest

- A randomized controlled basket trial with 1:1 randomization in 6 tumor indications, each targeting a hazard ratio of 0.5 in PFS with 90% power at 2.5% alpha
 - 88 PFS events and 110 patients planned for each indication
 - PFS analysis is conducted when all are enrolled
- D2 is applied to keep total sample size at 660 in pooled population targeting 430 death events
 - The study has ~90% power to detect a hazard ratio of 0.7 in OS at 0.8% alpha (after taking the penalty) assuming ρ =0.5
 - Observed hazard ratio ~0.79 or lower for a positive trial in pooled population (vs ~0.84 under D0)
- Potential to gain approvals in 6 indications based on comparable sample size to a conventional Phase 3 trial

Conclusions

- It is feasible to create a general design concept for a basket study that is suitable for many agents
- Multiple challenges can be addressed with careful planning
- Benefits include:
 - Increased and earlier patient access to targeted therapies for small subgroups
 - Cost-effective methods for sponsors to develop targeted agents in small subgroups
 - More robust datasets for health authorities to assess benefit-risk in these small patient groups

Key References

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